



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5 : A61K 48/00, 35/12, 39/00 C12N 15/19, 15/24, 15/25 C12N 15/26, 15/90, 15/63		A1 (11) International Publication Number: WO 93/07906 (43) International Publication Date: 29 April 1993 (29.04.93)
(21) International Application Number: PCT/US92/08999 (22) International Filing Date: 23 October 1992 (23.10.92)		(74) Agents: CAMPBELL, Cathryn et al.; Campbell & Flores, 4370 La Jolla Village Drive, Suite 700, San Diego, CA 92122 (US).
(30) Priority data: 781,356 25 October 1991 (25.10.91) US 863,641 3 April 1992 (03.04.92) US		(81) Designated States: CA, JP, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, SE).
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(54) Title: LYMPHOKINE GENE THERAPY OF CANCER		
(57) Abstract <p>A novel method of tumor immunotherapy is described comprising the genetic modification of cells resulting in the secretion of cytokine gene products to stimulate a patient's immune response to tumor antigens. In one embodiment, autologous fibroblasts genetically modified to secrete at least one cytokine gene product are utilized to immunize the patient in a formulation with tumor antigens at a site other than an active tumor site. In another embodiment, cells genetically modified to express at least one tumor antigen product and to secrete at least one cytokine gene product are utilized in a formulation to immunize the patient at a site other than an active tumor site.</p>		

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Lymphokine Gene Therapy of CancerBACKGROUND

This application is a continuation-in-part of United States Patent Application Serial No. 07/781,356, 5 filed on October 25, 1991, which is a continuation-in-part of United States Patent Application Serial No. 07/720,872, filed on June 25, 1991, both of which are incorporated herein in their entirety.

Recent advances in our understanding of the 10 biology of the immune system have lead to the identification of important modulators of immune responses, called cytokines (1-3). Immune system modulators produced by lymphocytes are termed lymphokines, a subset of the cytokines. These agents mediate many of the immune 15 responses involved in anti-tumor immunity. Several of these cytokines have been produced by recombinant DNA methodology and evaluated for their anti-tumor effects. The administration of lymphokines and related immunomodulators has resulted in objective tumor responses 20 in patients with various types of neoplasms (4-7). However, current modes of cytokine administration are frequently associated with toxicities that limit the therapeutic value of these agents.

For example, interleukin-2 (IL-2) is an important 25 lymphokine in the generation of anti-tumor immunity (4). In response to tumor antigens, a subset of lymphocytes termed helper T-cells secrete small quantities of IL-2. This IL-2 acts locally at the site of tumor antigen 30 stimulation to activate cytotoxic T-cells and natural killer cells which mediate systemic tumor cell destruction. Intravenous, intralymphatic and intralesional administration of IL-2 has resulted in clinically significant responses in some cancer patients (4-6). However, severe toxicities (hypotension and edema) limit 35 the dose and efficacy of intravenous and intralymphatic IL-

2 administration (5-7). The toxicity of systemically administered lymphokines is not surprising as these agents mediate local cellular interactions and they are normally secreted in only very small quantities.

5 Additionally, other cytokines, such as interleukin-4 (IL-4), alpha interferon (α -INF) and gamma interferon (γ -INF) have been used to stimulate immune responses to tumor cells. Like IL-2, the current modes of administration have adverse side effects.

10 To circumvent the toxicity of systemic cytokine administration, several investigators have examined intralesional injection of IL-2. This approach eliminates the toxicity associated with systemic IL-2 administration (8,9,10). However, multiple intralesional injections are 15 required to optimize therapeutic efficacy (9,10). Hence, these injections are impractical for many patients, particularly when tumor sites are not accessible for injection without potential morbidity.

20 An alternative approach, involving cytokine gene transfer into tumor cells, has resulted in significant anti-tumor immune responses in several animal tumor models (11-14). In these studies, the expression of cytokine gene products following cytokine gene transfer into tumor cells has abrogated the tumorigenicity of the cytokine-secreting 25 tumor cells when implanted into syngeneic hosts. The transfer of genes for IL-2 (11,12) γ -INF (13) or interleukin-4 (IL-4) (14) significantly reduced or eliminated the growth of several different histological types of murine tumors. In the studies employing IL-2 gene 30 transfer, the treated animals also developed systemic anti-tumor immunity and were protected against subsequent tumor challenges with the unmodified parental tumor (11,12). Similar inhibition of tumor growth and protective immunity was also demonstrated when immunizations were performed

with a mixture of unmodified parental tumor cells and genetically modified tumor cells engineered to express the IL-2 gene. No toxicity associates with localized lymphokine transgene expression was reported in these 5 animal tumor studies (11-14).

While the above gene-transfer procedure has been shown to provide anti-tumor immunity, it still retains practical difficulties. This approach is limited by the inability to transfer functional cytokine genes into many 10 patients' tumor cells, as most patients' tumors cannot be established to grow *in vitro* and methods for human *in vivo* gene transfer are not available.

SUMMARY OF THE INVENTION

The present invention demonstrates a novel, more 15 practical method of cytokine cancer immunotherapy. In one approach, selected cells from a patient, such as fibroblasts, obtained, for example, from a routine skin biopsy, are genetically modified to express one or more cytokines. Alternatively, patient cells which may normally 20 serve as antigen presenting cells in the immune system such as macrophages, monocytes, and lymphocytes may also be genetically modified to express one or more cytokines. These modified cells are hereafter called cytokine-expressing cells, or CE cells. The CE cells are then 25 mixed with the patient's tumor antigens, for example in the form of irradiated tumor cells, or alternatively in the form of purified natural or recombinant tumor antigen, and employed in immunizations, for example subcutaneously, to induce systemic anti-tumor immunity.

30 The cytokines are locally expressed at levels sufficient to induce or augment systemic anti-tumor immune responses via local immunization at sites other than active tumor sites. Systemic toxicity related to cytokine

administration should not occur because the levels of cytokine secreted by the CE cells should not significantly affect systemic cytokine concentrations.

As the amount of cytokine secreted by the CE 5 cells is sufficient to induce anti-tumor immunity but is too low to produce substantial systemic toxicity, this approach provides the benefit of local cytokine administration. In addition, this novel method obviates the need for intralesional injections, which may produce 10 morbidity. Furthermore, the continuous local expression of cytokine(s) at the sites of immunization may also augment anti-tumor immune responses compared to intermittent cytokine injections. This method also provides the advantage of local immunization with the CE cells, as 15 opposed to cumbersome intravenous infusions. This method also eliminates the need for establishing tumor cell lines in vitro as well as transfer of genes into these tumor cells.

This invention also provides an alternative means 20 of localized expression of cytokines to induce and/or increase immune responses to a patient's tumor through genetic modification of cellular expression of both cytokine(s) and tumor antigen(s). In this embodiment, selected cells from a patient are isolated and transduced 25 with cytokine gene(s) as well as gene(s) coding for tumor antigen(s). The transduced cells are called "carrier cells." Carrier cells can include fibroblasts and cells which may normally serve as antigen presenting cells in the immune system such as macrophages, monocytes, and 30 lymphocytes. Transduced carrier cells actively expressing both the cytokine(s) and the tumor antigen(s) are selected and utilized in local immunizations at a site other than active tumor sites to induce anti-tumor immune responses. As with the CE cells, these carrier cells should not 35 produce substantial systemic toxicities, as the levels of

cytokine(s) secreted by the carrier cells should not significantly affect systemic cytokine concentrations. This alternate embodiment is advantageous because it obviates the need to obtain samples of the tumor, which is 5 sometimes difficult. However, carrier cells can be utilized in local immunizations in conjunction with tumor cells, tumor cell homogenates, purified tumor antigens, or recombinant tumor antigens to enhance anti-tumor immunity.

Additionally, this second embodiment retains the 10 same advantages as the first embodiment in that the level of cytokine released by the carrier cells is sufficient to induce anti-tumor immunity but is too low to produce substantial systemic toxicity. In addition, as with the first embodiment, this method obviates the need for 15 intralesional injections, and allows for continuous expression of cytokine(s). This method also eliminates the need for establishing continuous cultures *in vitro* of tumor cells as well as transfer of genes into these tumor cells, and provides the advantage of local immunization with the 20 carrier cells, as opposed to cumbersome lengthy intravenous infusions.

These approaches may also find application in inducing or augmenting immune responses to other antigens of clinical significance in other areas of medical 25 practice.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows schematic diagrams of retroviral vectors DC/TKIL2, LXSN-IL2, and LNCX-IL2.

Figure 2 shows a mean IL-2 concentration of 30 triplicate supernatant samples measured by ELISA. Supernatants were harvested from overnight cultures of approximately 1.5×10^4 semi-confluent fibroblasts.

Figure 3 shows biological activity of the IL-2 secreted by the transduced fibroblasts was demonstrated by measuring mean ^3H -TdR incorporation of an IL-2 dependent T-cell line incubated with triplicate samples of supernatants. Supernatants were harvested from overnight cultures of approximately 1.5×10^6 semi-confluent fibroblasts.

Figure 4 shows comparisons between animals injected with 10^5 CT26 tumor cells alone (□); 10^5 CT26 tumor cells and 2×10^6 unmodified BALB/C fibroblasts (■); 10^5 CT26 tumor cells and 2×10^6 IL-2 transduced BALB/C fibroblasts (●); and 10^5 CT26 tumor cells and 1×10^6 transduced BALB/C fibroblasts (○). Tumor measurements are the mean products of the cross-sectional diameter of the tumors from four animals in each treatment group. The (*) indicates statistically significant difference ($P < 0.05$) in tumor growth curves.

Figure 5 shows PCR analysis of neomycin phosphotransferase DNA sequences. Lane 1 - positive control pLXSN-RI-IL2. Lanes 2 through 4 tests genomic DNA; Lanes 5 and 6 ovary genomic DNA; Lane 7 negative control, no DNA. Identical results were obtained with liver, spleen and lung genomic DNA (data not shown).

Figure 6 shows the effect of IL-2 modified fibroblasts on tumor establishment and development using 2×10^6 fibroblasts mixed with 5×10^6 CT26 tumor cells concentrating on the rate of tumor growth.

Figure 7 shows the effect of IL-2 modified fibroblasts on tumor establishment and development using 2×10^6 fibroblasts mixed with 5×10^6 CT26 tumor cells concentrating on the time of tumor onset for the individual animal in each treatment group.

Figure 8 shows the effect of IL-2 modified fibroblasts on tumor establishment and development using 2×10^4 fibroblasts mixed with 1×10^3 CT26 tumor cells concentrating on the rate of tumor growth.

5 Figure 9 shows the effect of IL-2 modified fibroblasts on tumor establishment and development using 2×10^4 fibroblasts mixed with 1×10^3 CT26 tumor cells concentrating on the time of tumor onset for the individual animal in each treatment group.

10 Figure 10 shows the effect of IL-2 modified cells on tumor establishment and development using 2×10^4 DCTK-IL2-modified CT26 tumor cells mixed with 1×10^3 unmodified CT26 compared to 2×10^4 DCTK-IL2-modified fibroblasts mixed with 1×10^3 CT26 concentrating on the rate of tumor growth.

15 Figure 11 shows the effect of IL-2 modified cells on tumor establishment and development using 2×10^4 DCTK-IL2-modified CT26 tumor cells mixed with 1×10^3 unmodified CT26 compared to 2×10^4 DCTK-IL2-modified fibroblasts mixed with 1×10^3 CT26 concentrating on the time of tumor onset 20 for the individual animal in each treatment group.

Figure 12 shows the effect of IL-2 modified fibroblasts on induction of systemic anti-tumor immunity and the rate of tumor growth. Mice were immunized with 2×10^4 fibroblasts mixed with 2.5×10^3 irradiated CT26 tumor 25 cells 7 days prior to challenge with 5×10^4 fresh tumor cells.

Figure 13 shows the effect of IL-2 modified fibroblasts on induction of systemic anti-tumor immunity and the time of tumor onset for the individual animal in 30 each treatment group. Mice were immunized with 2×10^4 fibroblasts mixed with 2.5×10^3 irradiated CT26 tumor cells 7 days prior to challenge with 5×10^4 fresh tumor cells.

Figure 14 shows the effect of IL-2 modified fibroblasts on induction of systemic anti-tumor immunity and the rate of tumor growth. Mice were immunized with 2×10^6 fibroblasts mixed with 2.5×10^5 irradiated CT26 tumor cells 14 days prior to challenge with 5×10^4 fresh tumor cells.

Figure 15 shows the effect of IL-2 modified fibroblasts on induction of systemic anti-tumor immunity and the time of tumor onset for the individual animal in each treatment group. Mice were immunized with 2×10^6 fibroblasts mixed with 2.5×10^5 irradiated CT26 tumor cells 14 days prior to challenge with 5×10^4 fresh tumor cells.

DETAILED DESCRIPTION

A novel method of tumor immunotherapy is described comprising the genetic modification of cells resulting in the secretion of cytokine gene products to stimulate a patient's immune response to tumor antigens. "Gene" is defined herein to be a nucleotide sequence encoding the desired protein. In one embodiment, autologous fibroblasts genetically modified to secrete at least one cytokine gene product are utilized to immunize the patient in a formulation with tumor antigens at a site other than an active tumor site. In another embodiment, cells genetically modified to express at least one tumor antigen gene product and to secrete at least one cytokine gene product are utilized in formulation to immunize the patient at a site other than an active tumor site. Cytokines are preferably expressed in cells which efficiently secrete these proteins into the surrounding milieu. fibroblasts are an example of such cells. Fibroblasts or other cells can be genetically modified to express and secrete one or more cytokines, as described later in this specification.

Tumor antigens can be provided by several methods, including, but not limited to the following: 1) CE cells can be transduced with gene(s) coding for tumor antigens. These "carrier cells" are then utilized in 5 patient immunizations. 2) Cloned gene sequences coding for appropriate tumor antigens can be transferred into cells such as fibroblasts or antigen-presenting cells. These cells are then mixed with CE or carrier cells to immunize the patient. 3) Tumor antigens can be cloned in bacteria 10 or other types of cells by recombinant procedures. These antigens are then purified and employed in immunization with CE and/or carrier cells. 4) Tumor antigens can be purified from tumor cells and used, along with CE or carrier cells, to immunize the patient. 5) Tumor cells may 15 be irradiated or mechanically disrupted and mixed with CE and/or carrier cells for patient immunizations.

This invention encompasses the following steps: (A) isolation of appropriate cells for generation of CE cells or carrier cells; (B) isolation of cytokine genes or 20 isolation of cytokine genes and tumor antigen genes, as well as appropriate marker and/or suicide genes; (C) transfer of the genes from (B) to produce the CE cells or carrier cells; (D) preparation of immunological samples of the patient's tumor antigens or other suitable tumor 25 antigens for immunization with CE or carrier cells; (E) inactivation of the malignant potential of tumor cells if they are used as a source of tumor antigens for immunization; and (F) preparation of samples for immunization. Following are several embodiments 30 contemplated by the inventors. However, it is understood that any means known by those in the art to accomplish these steps will be usable in this invention.

(A) Isolation of Cells to Generate CE and Carrier Cells

Cells to be utilized as CE cells and carrier cells can be selected from a variety of locations in the 5 patient's body. For example, skin punch biopsies provide a readily available source of fibroblasts for use in generating CE cells, with a minimal amount of intrusion to the patient. Alternatively, these fibroblasts can be obtained from the tumor sample itself. Cells of 10 hematopoietic origin may be obtained by venipuncture, bone marrow aspiration, lymph node biopsies, or from tumor samples. Other appropriate cells for the generation of CE or carrier cells can be isolated by means known in the art. Non-autologous cells similarly selected and processed can 15 also be used.

(B) Isolation of Genes

Numerous cytokine genes have been cloned and are available for use in this protocol. The genes for IL-2, γ -INF and other cytokines are readily available (1-5, 11-20 14). Cloned genes of the appropriate tumor antigens are isolated according to means known in the art.

Selectable marker genes such as neomycin 25 resistance (Neo^R) are readily available. Incorporation of a selectable marker gene(s) allows for the selection of cells that have successfully received and express the desired genes. Other selectable markers known to those in the art of gene transfer may also be utilized to generate CE cells or carrier cells expressing the desired transgenes.

30 "Suicide" genes can be incorporated into the CE cells or carrier cells to allow for selective inducible killing after stimulation of the immune response. A gene

such as the herpes simplex virus thymidine kinase gene (TK) can be used to create an inducible destruction of the CE cells or carrier cells. When the CE cells or carrier cells are no longer useful, a drug such as acyclovir or 5 gancyclovir can be administered. Either of these drugs will selectively kill cells expressing TK, thus eliminating the implanted transduced cells. Additionally, a suicide gene may be a gene coding for a non-secreted cytotoxic polypeptide attached to an inducible promoter. When 10 destruction of the CE or carrier cells is desired, the appropriate inducer of the promoter is administered so that the suicide gene is induced to produce cytotoxic polypeptide which subsequently kills the CE or carrier cell. However, destruction of the CE or carrier cells may 15 not be required.

Genes coding for tumor antigen(s) of interest can be cloned by recombinant methods. The coding sequence of an antigen expressed by multiple tumors may be utilized for many individual patients.

20 (C) Transfer of Genes

Numerous methods are available for transferring genes into cultured cells (15). For example, the appropriate genes can be inserted into vectors such as plasmids or retroviruses and transferred into the cells. 25 Electroporation, lipofection and a variety of other methods are known in the field and can be implemented.

One method for gene transfer is a method similar to that employed in previous human gene transfer studies, where tumor infiltrating lymphocytes (TILs) were modified 30 by retroviral gene transduction and administered to cancer patients (16). In this Phase I safety study of retroviral mediated gene transfer, TILs were genetically modified to express the Neomycin resistance (Neo^r) gene. Following

intravenous infusion, polymerase chain reaction analyses consistently found genetically modified cells in the circulation for as long as two months after administration. No infectious retroviruses were identified in these 5 patients and no side effects due to gene transfer were noted in any patients (16). These retroviral vectors have been altered to prevent viral replication by the deletion of viral gag, pol and env genes.

When retroviruses are used for gene transfer, 10 replication competent retroviruses may theoretically develop by recombination between the retroviral vector and viral gene sequences in the packaging cell line utilized to produce the retroviral vector. We will use packaging cell lines in which the production of replication competent 15 virus by recombination has been reduced or eliminated. Hence, all retroviral vector supernatants used to infect patient cells will be screened for replication competent virus by standard assays such as PCR and reverse transcriptase assays (16). Furthermore, exposure to 20 replication competent virus may not be harmful. In studies of subhuman primates injected with a large inoculum of replication competent murine retrovirus, the retrovirus was cleared by the primate immune system (17). No clinical 25 illnesses or sequelae resulting from replication competent virus have been observed three years after exposure. In summary, it is not expected that patients will be exposed to replication competent murine retrovirus and it appears that such exposure may not be deleterious (17).

30 (D) Preparation of Immunological Samples of the Patient's Tumor Antigens or Purified Recombinant Tumor Antigens

Tumor cells bearing tumor associated antigens are isolated from the patient. These cells can derive either from solid tumors or from leukemic tumors. For solid

tumors, single-cell suspensions can be made by mechanical separation and washing of biopsy tissue (18).

Hematopoietic tumors may be isolated from peripheral blood or bone marrow by standard methods (19).

5 A second variant is the use of homogenates of tumor cells. Such homogenates would contain tumor antigens available for recognition by the patient's immune system upon stimulation by this invention. Either unfractionated cell homogenates, made, for example, by mechanical 10 disruption or by freezing and thawing the cells, or fractions of homogenates preferably with concentrated levels of tumor antigens, can be used.

15 Likewise, purified tumor antigens, obtained for example by immunoprecipitation or recombinant DNA methods, could be used. Purified antigens would then be utilized for immunizations together with the CE cells and/or carrier cells described above to induce or enhance the patient's immune response to these antigens.

20 In the embodiments employing carrier cells, tumor antigens are available through their expression by the carrier cells. These carrier cells can be injected alone or in conjunction with other tumor antigen preparations or CE cells. Likewise, when CE cells are used, purified recombinant tumor antigen, produced by methods known in the 25 art, can be used.

If autologous tumor cells are not readily available, heterologous tumor cells, their homogenates, their purified antigens, or carrier cells expressing such antigens could be used.

(E) Inactivation of Tumor Cells

When viable tumor cells are utilized in immunizations as a source of tumor antigens, the tumor cells can be inactivated so that they do not grow in the patient. Inactivation can be accomplished by several methods. the cells can be irradiated prior to immunization (18). This irradiation will be at a level which will prevent their replication. Such viable calls can then present their tumor antigens to the patient's immune system, but cannot multiply to create new tumors.

Alternatively, tumor cells that can be cultured may be transduced with a suicide gene. As described above, a gene such as the herpes simplex thymidine kinase (TK) gene can be transferred into tumor cells to induce their destruction by administration of acyclovir or gancyclovir. After immunization, the TK expressing tumor cells can present their tumor antigens, and are capable of proliferation. After a period of time during which the patients's immune response is stimulated, the cells can be selectively killed. This approach might allow longer viability of the tumor cells utilized for immunizations, which may be advantageous in the induction or augmentation of anti-tumor immunity.

(F) Preparation of Samples for Immunization

CE cells and/or carrier cells and tumor cells, and/or homogenates of tumor cells and/or purified tumor antigen(s), are combined for patient immunization. Approximately 10^7 tumor cells will be required. If homogenates of tumor cells or purified or non-purified fractions of tumor antigens are used, the tumor dose can be adjusted based on the normal number of tumor antigens usually present on 10^7 intact tumor cells. The tumor preparation should be mixed with numbers of CE or carrier

cells sufficient to secrete cytokine levels that induce anti-tumor immunity (11-12) without producing substantial systemic toxicity which would interfere with therapy.

5 The cytokines should be produced by the CE cells or the carrier cells at levels sufficient to induce or augment immune response but low enough to avoid substantial systemic toxicity. This prevents side effects created by previous methods' administration of greater than physiological levels of the cytokines.

10 These mixtures, as well as carrier cells that are utilized alone, will be formulated for injection in any manner known in the art acceptable for immunization. Because it is important that at least the CE cells and carrier cells remain viable, the formulations must be 15 compatible with cell survival. Formulations can be injected subcutaneously, intramuscularly, or in any manner acceptable for immunization.

Contaminants in the preparation which may focus 20 the immune response on undesired antigens should be removed prior to the immunizations.

The following examples are provided for illustration of several embodiments of the invention and should not be interpreted as limiting the scope of the invention.

EXAMPLE IIMMUNIZATION WITH FIBROBLASTS EXPRESSING IL-2
MIXED WITH IRRADIATED TUMOR CELLS5 1) Isolation of Autologous Fibroblasts
for Use in Generating IL-2 Secreting CE Cells

Skin punch biopsies will be obtained from each patient under sterile conditions. The biopsy tissue will be minced and placed in RPMI 1640 media containing 10% fetal calf serum (or similar media) to establish growth of 10 the skin fibroblasts in culture. The cultured fibroblasts will be utilized to generate IL-2 secreting CE cells by retroviral mediated IL-2 gene transfer.

15 2) Retroviral Vector Preparation and
Generation of IL-2 Secreting CE Cells

18 The cultured skin fibroblasts will then be infected with a retroviral vector containing the IL-2 and Neomycin resistance (Neo^r) genes. An N2 vector containing the Neo^r gene will be used, and has been previously utilized by a number of investigators for in vitro and in vivo work, 20 including investigations with human subjects (16). The IL-2 vector will be generated from an N2-derived vector, LLRNL, developed and described by Friedmann and his colleagues (20). It will be made by replacement of the luciferase gene of LLRNL with a full-length cDNA encoding 25 human IL-2. Retroviral vector free of contaminating replication-competent virus is produced by transfection of vector plasmid constructions into the helper-free packaging cell line PA317. Before infection of patients' cells, the vector will have been shown to be free of helper virus. In 30 the event that helper virus is detected, the vector will be produced in the GP + envAM12 packaging cell line in which

the viral gag and pol genes are separated from the env, further reducing the likelihood of helper virus production.

3) Transduction Protocol

The cultured primary fibroblasts will be
5 incubated with supernatant from the packaging cell line as
described (20). Supernatant from these cells will be
tested for adventitious agents and replication competent
virus as described (16) and outlined in Table 1. The
fibroblasts are washed and then grown in culture media
10 containing G418, (a neomycin analogue) to select for
transduced cells expressing the Neo^r gene. The G418-
resistant cells will be tested for expression of the IL-2
gene by measuring the concentration of IL-2 in the culture
supernatant by an enzyme linked immunosorbent assay (ELISA)
15 (12). G418-resilient cells expressing IL-2 will be stored
at -70°C until required for subsequent use in
immunizations.

Table 1
Adventitious Agents and Safety Testing

20	1. Sterility
	2. Mycoplasma
	3. General Safety
	4. Viral Testing
	LCM Virus
25	Thymic agent
	S+/L- eco
	S+/L-xeno
	S+/L- ampho
	3T3 amplification
30	MRC-5/Vero

4) Preparation of Irradiated Tumor Cells

Tumors obtained from clinically indicated surgical resections or from superficial lymph node or skin metastases will be minced into 2-3 mm pieces and treated 5 with collagenase and DNase to facilitate separation of the tumor into a single cell suspension. The collected cells will be centrifuged and washed in RPMI 1640 media and then cryopreserved in a solution containing 10% dimethyl sulphoxide and 50% fetal calf serum in RPMI 1640 media. 10 The cells will be stored in liquid nitrogen until the time of administration. Prior to their use in subcutaneous immunizations, the cells will be thawed, washed in media free of immunogenic contaminants, and irradiated with 4,000 rads per minute for a total of 20,000 rads in a cesium 15 irradiator.

5) Patient Selection

Patients will have a histologically confirmed diagnosis of cancer. Patients with tumors that must be resected for therapeutic purposes or with tumors readily 20 accessible for biopsy are most appropriate for this embodiment of the invention.

6) Pretreatment Evaluation

The following pretreatment evaluations will be performed:

25 1) History and physical examination including a description and quantification of disease activity.

2) Performance Status Assessment

5
0 = Normal, no symptoms
1 = Restricted, but ambulatory
2 = Up greater than 50% of waking hours, capable of self-care
3 = Greater than 50% of waking hours confined to bed or chair, limited self-care
4 = Bedridden

10 3) Pretreatment Laboratory:

CBC with differential, platelet count, PT, PTT, glucose, BUN, creatinine, electrolytes, SGOT, SGPT, LDH, alkaline phosphatase, bilirubin, uric acid, calcium, total protein albumin.

15 4) Other Analyses:

Urinalysis

CH₅₀, C₁ and C₃ serum complement levels
Immunophenotyping of peripheral blood B cell and T cell subsets

20 Assays for detectable replication-competent virus in peripheral blood cells

PCR assays of peripheral blood leukocytes for Neo^A, IL-2 and viral env

5) Other Pretreatment Evaluation:

25 Chest X-ray and other diagnostic studies including computerized tomography (CT), magnetic resonance imaging (MRI) or radionuclide scans may be performed to document and quantify the extent of disease activity.

30 Follow-up evaluations of these assessments at regular intervals during the course of therapy (approximately every 1 to 3 months) will be useful in determining response to therapy and potential toxicity,

permitting adjustments in the number of immunizations administered.

7) Restrictions on Concurrent Therapy

For optimal effects of this treatment, patients 5 should receive no concurrent therapy which is known to suppress the immune system.

8) Final Formulation

Each patient will receive subcutaneous 10 immunizations with a mixture of irradiated tumor cells and autologous fibroblast CE cells genetically modified to secrete IL-2. Approximately 10^7 tumor cells will be mixed with 10^7 fibroblasts known to secrete at least 20 units/ml of IL-2 in tissue culture when semi-confluent (12). The 15 irradiated tumor cells and genetically modified fibroblasts will be placed in a final volume of 0.2 ml normal saline for immunization.

9) Dose Adjustments

At least two subcutaneous immunizations will be administered, two weeks apart, with irradiated tumor cells 20 and autologous fibroblasts genetically modified to secrete IL-2. If no toxicity is observed, subsequent booster immunizations may be administered periodically (at least one week apart) to optimize the anti-tumor immune response.

J) Treatment of Potential Toxicity

25 Toxic side effects are not expected to result from these immunizations. However, potential side effects of these immunizations are treatable in the following manner:

If massive tumor cell lysis results, any resulting uric acid nephropathy, adult respiratory distress syndrome, disseminated intravascular coagulation or hyperkalemia will be treated using standard methods.

5 Local toxicity at the sites of immunization will be treated with either topical steroids and/or surgical excision of the injection site as deemed appropriate.

10 Hypersensitivity reactions such as chills, fever and/or rash will be treated symptomatically with antipyretics and antihistamines. Patients should not be treated prophylactically. Should arthralgias, 15 lymphadenopathy or renal dysfunction occur, treatment with corticosteroids and/or antihistamines will be instituted. Anaphylaxis will be treated by standard means such as administration of epinephrine, fluids, and steroids.

EXAMPLE II

A. Retroviral IL-2 Gene Transfer and Expression in Fibroblasts

20 Retroviral vectors were employed to transfer and express IL-2 and neomycin phosphotransferase genes in murine and primary human fibroblasts. The retroviral vector DC/TKIL2 produced by Gilboa and co-workers (Gansbacher, et al., J. Exp. Med. 172:1217-1223, 1990, which is incorporated herein by reference) was utilized to 25 transduce murine fibroblasts for application in an animal tumor model (see Section B below). Human fibroblasts were transduced with the retroviral vector LXSN-RI-IL2. Schematic diagrams of the structure of these retroviral 30 vectors are provided in Figure 1. A more complete description of the LXSN-RI-IL2 vector, including its nucleotide sequence, is provided in Example III and in Tables 2, 3 and 4.

Following infection with the described vectors and selection for 2-3 weeks in growth media containing the neomycin analogue G418, Balb/c and human embryonic fibroblast culture supernatants were harvested and tested 5 for IL-2 by an enzyme-linked immunosorbent assay (ELISA). Figure 2 depicts the levels of IL-2 secreted by the transduced fibroblasts.

These results can be confirmed using negative control fibroblasts infected with an N2-derived retroviral 10 vector expressing an irrelevant gene such as luciferase or β -galactosidase and studies with adult human fibroblasts.

Biological activity of the IL-2 expressed by the transduced human fibroblasts was confirmed by a cell proliferation bioassay employing an IL-2 dependent T cell 15 line. In this assay, supernatant from the transduced fibroblasts and control unmodified fibroblasts were incubated with the IL-2 dependent T cell line CTLL-2. Incorporation of ^3H -thymidine was measured as an indicator of cell proliferation and IL-2 activity (Figure 3).

20 B. Efficacy of Transduced Fibroblasts in an Animal Tumor Model

The efficacy of fibroblasts genetically modified to secrete IL-2 was tested in an animal model of colorectal carcinoma. In these studies, the Balb/c CT26 tumor cell 25 line was injected subcutaneously with Balb/c fibroblasts transduced to express IL-2. Control groups included animals injected with 1) a mixture of CT26 tumor cells and unmodified fibroblasts; 2) CT26 tumor cells without fibroblasts and 3) transduced fibroblasts alone. No tumors 30 were detected in 3/8 animals treated with transduced fibroblasts and CT26 cells. In contrast, all untreated control animals (8/8) injected with CT26 tumor cells developed palpable tumors. No tumors were detected in the

5 animals inoculated with transduced fibroblasts without CT26 tumor cells. The mean CT26 tumor size in Balb/c mice injected with the IL-2 secreting fibroblasts was considerably smaller compared to the control groups (Figure 4). A multivariate non-parametric statistical procedure (Koziol, et al., Biometrics 37:383-390, 1981 and Koziol, et al., Computer Prog. Biomed. 19:69-74, 1984, which is incorporated herein by reference) was utilized to evaluate differences in tumor growth among the treatment groups.

10 10 The tumor growth curves for the four treatment groups presented in Figure 4 were significantly different (p=0.048). Subsequent comparisons between treatment groups revealed a significant difference (p < 0.05) in tumor growth between animals injected with CT26 tumor cells alone

15 15 and animals treated with 2×10^6 transduced fibroblasts and CT26 tumor cells (Figure 4).

EXAMPLE III

A. Project Overview

20 20 Lymphokine gene therapy of cancer will be evaluated in cancer patients who have failed conventional therapy. An N2-derived vector containing the neomycin phosphotransferase gene will be used. This vector has been employed by a number of investigators for in vitro and in vivo studies including recently approved investigations 25 25 with human subjects (Rosenberg et al., N. Eng. J. Med., 323:570-578, 1990). The lymphokine vectors used in this investigation will be generated from the N2-derived vector, LXSN, developed and described by Miller et al., Mol. Cell Biol. 6:2895, 1986 and Miller et al., BioTechniques 7:980, 30 30 1989, which are incorporated herein by reference. The vector LXSN-RI-IL2 contains human IL-2 cDNA under the control of the retroviral 5' LTR promoter and the neomycin phosphotransferase gene under the control of the SV40 promoter (see Figure 1). The normal human IL-2 leader

sequence has been replaced with a chimeric sequence containing rat insulin and human IL-2 leader sequences (see Tables 2, 3 and 4). This chimeric leader sequence enhances IL-2 gene expression.

5 To construct the LXSN-RI-IL2 vector, the bacterial plasmid pBC12/CMV/IL2 (Cullen, B.R., DNA 7:645-650, 1988, which is incorporated herein by reference) containing the full-length IL-2 cDNA and chimeric leader sequence was digested with HindIII and the ends were
10 blunted using Klenow polymerase. IL-2 cDNA was subsequently released from the plasmid by digestion with BamHI. The IL-2 fragment was purified by electrophoresis in a 1% agarose gel and the appropriate band was extracted utilizing a glass powder method. Briefly, the gel slice
15 was dissolved in 4M NaI at 55°. After cooling to room temperature, 4 μ l of oxidized silica solution (BIO-101, La Jolla, CA) was added to adsorb the DNA. The silica was then washed with a cold solution of 50% ethanol containing 0.1 M NaCl in TE buffer. The DNA was eluted from the
20 silica by heating at 55° in distilled H₂O. The purified IL-2 cDNA was then directionally ligated into the HpaI-BamHI cloning sites of the pLXSN vector. A more complete description of the pLXSN-RI-IL2 vector and its partial
25 nucleotide sequence are provided in Tables 2, 3, 4, 5 and 6.

Table 2

Description of the LXSN-RJ-IL2
from position 1 to 6365

<u>Bases</u>	<u>Description</u>
1-589	Moloney murine sarcoma virus 5' LTR
659-1458	The sequence of the extended packaging signal
1469-2151	IL-2 cDNA with chimeric leader sequence
1469-1718	IL-2 chimeric leader sequence
1647-1718	coding region of the signal peptide
1719-2151	Mature IL-2 coding sequence
2158-2159	Mo mu sarcoma virus end/SV 40 start
2159-2503	Simian virus 40 early promoter
2521-2522	Simian virus DNA end/Tn5 DNA start
2557-3351	Neomycin phosphotransferase
3370-3371	Tn5 DNA end/Moloney murine leukemia virus start
3411-4004	Moloney murine leukemia virus 3' LTR
4073-4074	Moloney murine leukemia DNA end/pBR322 DNA start
4074-6365	Plasmid backbone

Table 3

Enzyme	[# Cuts	Position(s)
Aat1	[2]	1961, 2481
Aat2	[2]	811, 6295
Acc1	[1]	4252
Acc2	[19]	392, 394, 445, 969, 971, 1193, 2751, 3052, 3084, 3807, 3809, 4081, 4083, 4186, 4527, 5108, 5438, 5931, 6263
Acy1	[5]	808, 2685, 3860, 5910, 6292
Afl1	[13]	260, 273, 328, 626, 756, 1277, 3201, 3676, 3689, 3744, 4041, 5511, 5733
Afl2	[4]	34, 1064, 1955, 3446
Afl3	[2]	1592, 4480
Ahal	[20]	161, 237, 473, 474, 602, 644, 789, 2689, 2849, 3578, 3653, 3888, 3889, 4017, 4059, 4126, 4161, 4860, 5556, 5907
Aha2	[5]	808, 2685, 3860, 5910, 6292
Aha3	[3]	5239, 5258, 5950
Alul	[33]	29, 33, 119, 190, 411, 654, 734, 742, 1470, 1486, 1751, 1935, 2003, 2446, 2500, 2791, 3249, 3441, 3445, 3532, 3607, 3826, 4069, 4122, 4141, 4422, 4648, 4738, 4784, 5041, 5562, 5662, 5725
Alw1	[20]	1110, 1414, 1665, 2018, 2147, 2160, 2529, 2553, 2864, 2929, 3110, 4027, 5041, 5127, 5129, 5225, 5226, 5689, 6006, 6010
AlwN1	[4]	231, 3572, 3647, 4896
Aoc1	[2]	847, 1076
Aoc2	[19]	323, 413, 426, 597, 1583, 1721, 2631, 2724, 2798, 2988, 3050, 3739, 3828, 3841, 4012, 4300, 4798, 5959, 6044
Aos1	[2]	2787, 5595
Apal1	[4]	1717, 4296, 4794, 6040

Apy1	[22]	315, 623, 801, 814, 1227, 1252, 1275, 1295, 1325, 1526, 1536, 1558, 1630, 2196, 2251, 2268, 3072, 3731, 4038, 4508, 4629, 4642
Aql1	[6]	241, 472, 1998, 3821, 3854, 3887
Asel	[2]	1801, 5545
Asp700	[1]	5972
Asp718	[2]	476, 3891
AspA1	[1]	1145
Asul	[29]	169, 200, 245, 260, 273, 328, 626, 756, 826, 839, 1043, 1254, 1277, 1532, 1649, 3201, 3541, 3586, 3616, 3661, 3676, 3689, 3744, 4041, 5415, 5494, 5511, 5733, 6349
Aval	[6]	241, 472, 1998, 3821, 3854, 3887
Ava2	[13]	260, 273, 328, 626, 756, 1277, 3201, 3676, 3689, 3744, 4041, 5511, 5733
Ava3	[2]	2232, 2304
Avr2	[2]	1962, 2482
Ball	[3]	658, 1169, 2767
BamH1	[1]	2152
Ban1	[9]	318, 476, 1200, 2684, 2719, 3734, 3859, 3891, 5321
Ban2	[8]	413, 426, 597, 1583, 3050, 3828, 3841, 4012
Bbel	[2]	2688, 3863
Bbv1	[22]	969, 997, 1738, 2493, 2632, 2758, 2800, 2816, 2909, 3321, 4060, 4131, 4228, 4372, 4390, 4809, 4899, 4902, 5108, 5411, 5600, 5802
Bcl1	[1]	2526
Bgl1	[2]	2435, 5493
Bsp1286I	[19]	323, 413, 426, 597, 1583, 1721, 2631, 2724, 2798, 2988, 3050, 3739, 3828, 3841, 4012, 4300, 4798, 5959, 6044

BspH1	[3]	5200,	6208,	6313				
BspM1	[4]	1501,	2500,	2572,	2953			
BssH2	[4]	392,	443,	3082,	3807			
BstE2	[1]	1145						
BstN1	[22]	315,	623,	801,	814,	1227,	1252,	
	1275,	1295,	1325,	1526,	1536,	1558,	1630,	
	2196,	2251,	2268,	3072,	3731,	4038,	4508,	
	4629,	4642						
BstU1	[19]	392,	394,	445,	969,	971,	1193,	
	2751,	3052,	3084,	3807,	3809,	4081,	4083,	
	4186,	4527,	5108,	5438,	5931,	6263		
BstX1	[1]	2060						
BstY1	[11]	2010,	2152,	2521,	2856,	3102,	5121,	
	5132,	5218,	5230,	5998,	6015			
Bsu36I	[2]	847,	1076					
Ccr1	[1]	1998						
Cfol	[31]	394,	396,	445,	447,	714,	971,	
	2679,	2687,	2751,	2788,	3054,	3084,	3086,	
	3314,	3809,	3811,	3862,	4083,	4186,	4216,	
	4357,	4390,	4660,	4727,	4827,	5001,	5110,	
	5503,	5596,	5933,	6265				
Cfr1	[9]	656,	790,	1167,	1188,	2591,	2765,	
	3156,	3183,	5761					
Cfr10I	[3]	3004,	3185,	5453				
Cfr13I	[29]	169,	200,	245,	260,	273,	328,	
	626,	756,	826,	839,	1043,	1254,	1277,	
	1532,	1649,	3201,	3541,	3586,	3616,	3661,	
	3676,	3689,	3744,	4041,	5415,	5494,	5511,	
	5733,	6349						
Cvn1	[2]	847,	1076					
Ddel	[23]	75,	165,	191,	282,	553,	847,	
	1076,	1348,	1692,	2442,	3348,	3487,	3582,	
	3657,	3698,	3879,	3967,	4290,	4755,	5164,	
	5330,	5870,	6296					
Dpn1	[30]	95,	1104,	1236,	1421,	1659,	2012,	
	2154,	2523,	2528,	2547,	2858,	2936,	3017,	
	3026,	3104,	3507,	4021,	5048,	5123,	5134,	
	5142,	5220,	5232,	5337,	5678,	5696,	5742,	
	6000,	6017,	6053					

29

Dral	[3]	5239,	5258,	5950			
Dra2	[4]	328,	1277,	3744,	6349		
Eael	[9]	656,	790,	1167,	1188,	2591,	2765,
	3156,	3183,	5761				
Eag1	[2]	790,	2591				
Eco47I	[13]	260,	273,	328,	626,	756,	1277,
	3201,	3676,	3689,	3744,	4041,	5511,	5733
Eco52I	[2]	790,	2591				
Eco81I	[2]	847,	1076				
EcoN1	[2]	850,	1450				
EcoO109I	[4]	328,	1277,	3744,	6349		
EcoR1	[1]	1460					
EcoR1*	[14]	938,	1037,	1460,	1798,	1805,	1928,
	2064,	2121,	2236,	2308,	2400,	5240,	5546,
	5801						
EcoR2	[22]	313,	621,	799,	812,	1225,	1250,
	1273,	1293,	1323,	1524,	1534,	1556,	1628,
	2194,	2249,	2266,	3070,	3729,	4036,	4506,
	4627,	4640					
EcoR5	[4]	137,	213,	3554,	3629		
EcoT22I	[2]	2232,	2304				
Fdi2	[2]	2787,	5595				
Fnu4H1	[41]	793,	967,	983,	986,	1191,	1752,
	2430,	2507,	2594,	2646,	2657,	2747,	2752,
	2789,	2830,	2917,	2920,	2923,	3159,	3255,
	3296,	3310,	4074,	4120,	4217,	4270,	4386,
	4404,	4407,	4525,	4680,	4823,	4888,	4891,
	5097,	5425,	5614,	5764,	5791,	5886,	6115
FnuD2	[19]	392,	394,	445,	969,	971,	1193,
	2751,	3052,	3084,	3807,	3809,	4081,	4083,
	4186,	4527,	5108,	5438,	5931,	6263	
Fok1	[13]	498,	1198,	1358,	1679,	2333,	2552,
	3009,	3034,	3912,	4168,	5339,	5520,	5807
Fsp1	[2]	2787,	5595				
Hae2	[4]	2688,	3863,	4358,	4728		

Hae3	[35]	171, 202, 247, 658, 792, 828,
	840, 1045, 1169, 1190, 1255, 1534, 1650,	
	1866, 1961, 2423, 2429, 2438, 2481, 2593,	
	2767, 3158, 3185, 3543, 3588, 3618, 3663,	
	4495, 4506, 4524, 4958, 5416, 5496, 5763,	
	6350	
Hpa2	[30]	161, 237, 473, 601, 643, 789,
	2590, 2667, 2689, 2717, 2848, 2938, 3005,	
	3186, 3578, 3653, 3888, 4016, 4058, 4126,	
	4160, 4687, 4834, 4860, 5050, 5454, 5488,	
	5555, 5665, 5907	
Bgl1	[8]	455, 707, 960, 1580, 4175, 4591,
	5169, 5899	
BgiA1	[9]	413, 1721, 2798, 2988, 3828, 4300,
	4798, 5959, 6044	
Bhal	[31]	394, 396, 445, 447, 714, 971,
	2679, 2687, 2751, 2788, 3054, 3084, 3086,	
	3314, 3809, 3811, 3862, 4083, 4186, 4216,	
	4357, 4390, 4660, 4727, 4827, 5001, 5110,	
	5503, 5596, 5933, 6265	
HinP1	[31]	392, 394, 443, 445, 712, 969,
	2677, 2685, 2749, 2786, 3052, 3082, 3084,	
	3312, 3807, 3809, 3860, 4081, 4184, 4214,	
	4355, 4388, 4658, 4725, 4825, 4999, 5108,	
	5501, 5594, 5931, 6263	
Hinc2	[1]	5914
Hind2	[1]	5914
Hind3	[1]	2498
Hinf1	[14]	298, 517, 857, 868, 1553, 1814,
	3170, 3304, 3356, 3881, 4380, 4455, 4851,	
	5368	
Hpa2	[30]	161, 237, 473, 601, 643, 789,
	2590, 2667, 2689, 2717, 2848, 2938, 3005,	
	3186, 3578, 3653, 3888, 4016, 4058, 4126,	
	4160, 4687, 4834, 4860, 5050, 5454, 5488,	
	5555, 5665, 5907	
Hph1	[11]	1214, 1240, 1817, 2863, 4102, 4111,
	5216, 5443, 5859, 6065, 6100	
Kpn1	[2]	480, 3895
Mae1	[15]	30, 293, 689, 727, 739, 1452,
	1606, 1893, 1963, 2483, 3442, 3709, 4975,	
	5228, 5563	

Mae2	[11]	808, 1139, 1180, 1987, 2801, 2988, 4233, 5183, 5599, 5972, 6292
Mae3	[20]	38, 1052, 1080, 1145, 1289, 1478, 1706, 2805, 3111, 3450, 4134, 4229, 4836, 4899, 5015, 5298, 5629, 5687, 5840, 6028
Mbo1	[30]	93, 1102, 1234, 1419, 1657, 2010, 2152, 2521, 2526, 2545, 2856, 2934, 3015, 3024, 3102, 3505, 4019, 5046, 5121, 5132, 5140, 5218, 5230, 5335, 5676, 5694, 5740, 5998, 6015, 6051
Mbo2	[17]	444, 1145, 1356, 1575, 1617, 1908, 1911, 3046, 3256, 3336, 4351, 5142, 5213, 5968, 6046, 6155, 6351
Mnl1	[54]	291, 444, 508, 534, 560, 639, 841, 939, 1227, 1330, 1363, 1369, 1372, 1378, 1408, 1411, 1426, 1433, 1449, 1559, 1620, 1909, 1921, 2412, 2418, 2443, 2449, 2455, 2458, 2470, 2508, 2535, 2599, 2735, 3092, 3286, 3707, 3859, 3878, 3923, 3948, 3974, 4054, 4087, 4117, 4379, 4587, 4662, 4911, 5311, 5392, 5540, 5746, 6339
Msel	[22]	35, 1065, 1177, 1207, 1231, 1801, 1843, 1956, 1971, 2124, 2139, 3447, 4261, 5186, 5238, 5243, 5257, 5310, 5545, 5584, 5949, 6321
Msp1	[30]	161, 237, 473, 601, 643, 789, 2590, 2667, 2689, 2717, 2848, 2938, 3005, 3186, 3578, 3653, 3888, 4016, 4058, 4126, 4160, 4687, 4834, 4860, 5050, 5454, 5488, 5555, 5665, 5907
Mst1	[2]	2787, 5595
Mst2	[2]	847, 1076
Mval	[22]	315, 623, 801, 814, 1227, 1252, 1275, 1295, 1325, 1526, 1536, 1558, 1630, 2196, 2251, 2268, 3072, 3731, 4038, 4508, 4629, 4642
Nael	[1]	3187
Nar1	[2]	2685, 3860
Nc11	[20]	161, 237, 473, 474, 602, 644, 789, 2689, 2849, 3578, 3653, 3888, 3889, 4017, 4059, 4126, 4161, 4860, 5556, 5907
Ncol	[2]	2389, 3117

Nde1	[1]	4303
Nde2	[30]	93, 1102, 1234, 1419, 1657, 2010, 2152, 2521, 2526, 2545, 2856, 2934, 3015, 3024, 3102, 3505, 4019, 5046, 5121, 5132, 5140, 5218, 5230, 5335, 5676, 5694, 5740, 5998, 6015, 6051
Nhe1	[3]	29, 1605, 3441
Nla3	[26]	61, 1263, 1596, 1649, 1835, 1856, 2030, 2230, 2302, 2393, 2559, 2904, 3090, 3121, 3147, 3473, 4119, 4224, 4484, 5204, 5695, 5705, 5783, 5819, 6212, 6317
Nla4	[28]	153, 246, 262, 320, 478, 627, 758, 827, 959, 1202, 1279, 2154, 2200, 2272, 2686, 2721, 3678, 3736, 3861, 3893, 4042, 4512, 4551, 5323, 5417, 5458, 5669, 6259
Nsil	[2]	2232, 2304
Nsp(7524)1	[8]	1596, 1835, 1856, 2230, 2302, 3090, 4119, 4484
Nsp(7524)2	[19]	323, 413, 426, 597, 1583, 1721, 2631, 2724, 2798, 2988, 3050, 3739, 3828, 3841, 4012, 4300, 4798, 5959, 6044
NspB2	[12]	119, 190, 1751, 2158, 2791, 3532, 3607, 3989, 4192, 4822, 5067, 6008
NspH1	[8]	1596, 1835, 1856, 2230, 2302, 3090, 4119, 4484
PaeR7I	[1]	1998
Pall	[35]	171, 202, 247, 658, 792, 828, 840, 1045, 1169, 1190, 1255, 1534, 1650, 1866, 1961, 2423, 2429, 2438, 2481, 2593, 2767, 3158, 3185, 3543, 3588, 3618, 3663, 4495, 4506, 4524, 4958, 5416, 5496, 5763, 6350
PleI	[7]	865, 1547, 3350, 3889, 4374, 4859, 5362
PpuM1	[3]	328, 1277, 3744
Pss1	[4]	331, 1280, 3747, 6352
Pst1	[6]	987, 1163, 1888, 2511, 2738, 5618
PvuI	[1]	5743

Pvu2	[6]	119,	190,	1751,	2791,	3532,	3607
Rsal	[10]	347,	478,	725,	1342,	1519,	1597,
	2991,	3893,	4288,	5853			
Rsr2	[1]	3201					
Sac1	[2]	413,	3828				
Saul	[2]	847,	1076				
Sau3Al	[30]	93,	1102,	1234,	1419,	1657,	2010,
	2152,	2521,	2526,	2545,	2856,	2934,	3015,
	3024,	3102,	3505,	4019,	5046,	5121,	5132,
	5140,	5218,	5230,	5335,	5676,	5694,	5740,
	5998,	6015,	6051				
Sau96I	[29]	169,	200,	245,	260,	273,	328,
	626,	756,	826,	839,	1043,	1254,	1277,
	1532,	1649,	3201,	3541,	3586,	3616,	3661,
	3676,	3689,	3744,	4041,	5415,	5494,	5511,
	5733,	6349					
Scal	[1]	5853					
Scrf1	[42]	161,	237,	315,	473,	474,	602,
	623,	644,	789,	801,	814,	1227,	1252,
	1295,	1325,	1526,	1536,	1558,	1630,	2196,
	2251,	2268,	2689,	2849,	3072,	3578,	3653,
	3731,	3888,	3889,	4017,	4038,	4059,	4126,
	4161,	4508,	4629,	4642,	4860,	5556,	5907
Sdul	[19]	323,	413,	426,	597,	1583,	1721,
	2631,	2724,	2798,	2988,	3050,	3739,	3828,
	3841,	4012,	4300,	4798,	5959,	6044	
Sec1	[38]	159,	235,	314,	324,	472,	536,
	621,	622,	760,	799,	800,	812,	813,
	1294,	1303,	1323,	1324,	1525,	1557,	1962,
	2194,	2266,	2389,	2424,	2433,	2482,	2848,
	3117,	3576,	3651,	3730,	3740,	3887,	3950,
	4036,	4037,	4640				
SfaN1	[23]	258,	520,	997,	1657,	2107,	2239,
	2311,	2643,	2898,	2984,	3048,	3114,	3323,
	3674,	3934,	4146,	4281,	4317,	4357,	4577,
	5629,	5820,	6069				
Sfil	[1]	2435					
Smal	[2]	474,	3889				
Spel	[1]	726					
Sph1	[4]	1835,	2230,	2302,	3090		

Ssp1	[1]	6177
Sst1	[2]	413, 3828
Stu1	[2]	1961, 2481
Sty1	[9]	324, 536, 1303, 1962, 2389, 2482 , 3117, 3740, 3950
Taq1	[15]	860, 1096, 1407, 1418, 1660, 1999, 2514, 2798, 2954, 2978, 3014, 3176, 3367, 4580, 6024
Tha1	[19]	392, 394, 445, 969, 971, 1193, 2751, 3052, 3084, 3807, 3809, 4081, 4083, 4186, 4527, 5108, 5438, 5931, 6263
Tth111I	[6]	465, 877, 1275, 2803, 3880, 4227
Xba1	[2]	1892, 3708
Xho1	[1]	1998
Xho2	[11]	2010, 2152, 2521, 2856, 3102, 5121, 5132, 5218, 5230, 5998, 6015
Xma1	[2]	472, 3887
Xma3	[2]	790, 2591
Xmn1	[1]	5972
Xor2	[1]	5743

Table 4

Enzymes which do not cut LXSNRII.L2:

Acc3	Bgl2	Cla1	Hpa1	Nru1
SnaB1				
Apal	Bsm1	Dra3	Mlu1	PflM1
Sph1				
Asu2	BspM2	Eco47III	Mro1	Sac2
Sst2				
Ban3	BstB1	Esp1	Not1	Sall

Table 5

From 1 to 6365. Numbered from position 1.

	LXSNR11.L2	1000+	2000+	3000+	4000+	5000+	6000+
No-MuSV 5' long ter	-	-	-	-	-	-	-
- [Split]	-	-	-	-	-	-	-
1 to 683 of RIL2	>	>	>	>	>	>	>
neomycin phosphotra							
No-MuLV 3' long ter							
signal	1	+	1	1	1	1	1
Aat1	-	-	-	-	-	-	-
Aat2	-	1	+	+	+	+	+
Acc1	-	-	+	+	+	+	+
Acc2	-	-	-	-	-	-	-
Acy1	-	-	-	-	-	-	-
Afl1	-	-	-	-	-	-	-
Afl1	-	-	-	-	-	-	-
Afl2	-	-	-	-	-	-	-
Afl3	-	-	-	-	-	-	-
Aha1	-	-	-	-	-	-	-
Aha2	-	-	-	-	-	-	-
Aha3	-	-	-	-	-	-	-
Alu1	2	1	-12	-2	-2	-1	-1
Alu1	-	-	-	-	-	-	-
Alw1	-	-	-	-	-	-	-
Alw1	-	-	-	-	-	-	-
Aoc1	-	-	-	-	-	-	-
Aoc2	-	-	-	-	-	-	-
Aos1	-	-	-	-	-	-	-
Apal1	-	-	-	-	-	-	-
Ap1	-	-	-1-2-+23-31	-3	-1	-1-2	-1
Aqu1	-	-	-	-	-	-	-
Asel1	-	-	-	-	-	-	-
Asp100	-	-	-	-	-	-	-
Asp118	-	-	-	-	-	-	-
Asp1	-	-	-	-	-	-	-

SUBSTITUTE SHEET

EcoB11	-+1-	+-----
EcoB1	-+1-	-1-----
EcoB1091	-+1-	-1-----
EcoB1	-+1-	-1-----
EcoB1*	-+1-	-1-----
EcoB2	-+1-	-21+11111-----
EcoB2	-+1-	-23-31-----+3-----
EcoB5	-+1-	-+-----
EcoP221	-+1-	-+-----
Fdi2	-+1-	-+-----
FnuH1	-+1-	-111223+112-----
FnuH2	-+1-	-2+1-----
Fok1	-+1-	-1-----+2-----
Fsp1	-+1-	-1-----+1-----
Hae2	-+1-	-+-----
Hae3	-+1-	-3+1122-11-11-----
Hae3	-+1-	-221-1-----+2-----
Hep2	-+1-	-12111-1-----+2-----
Hga1	-+1-	-1-----+4-----
Hgl11	-+1-	-1-----+1-----
Hha1	-+1-	-211+3-----1-----
HinP1	-+1-	-211+3-----1-----
Hinc2	-+1-	-+-----
Hind2	-+1-	-+-----
Hind3	-+1-	-1-----
HinP1	-+1-	-2+-----
Hpa2	-+1-	-1-----+1-----
Hph1	-+1-	-2+-----
Kpn1	-+1-	-+-----
Mae1	-+1-	-3+-----1-----
Mae1	-+1-	-1-----+1-----
Mae2	-+1-	-1-----+1-----
Mae3	-+1-	-21+1-1-----+1-----
Mbo1	-+1-	-1-----+1-----
Mbo2	-+1-	-117211-----2+-----
Mnl1	-+1-	-131-11-----1-----
Mse1	-+1-	-3+1-2-----+2-----
Msp1	-+1-	-112111-1-----+1-----
Msp1	-+1-	-1-----+1-----
Msp2	-+1-	-1-----+1-----

Sph1
Ssp1
Sst1
Stu1
Sty1
Taq1
Thal
Tth1111
Xba1
Xho1
Xho2
Xma1
Xma3
Xmn1
Xor2

Table 5

from 1 to 655. Numbered from position 1.

Figure 2: Electropherogram of the 603 of RIL3. The plot shows signal intensity (y-axis, 0 to 6000) versus time (x-axis, 0 to 10000). The signal is labeled "603 of RIL3" and "Heavy 2, long tail". The plot includes a legend for "Acetyl phosphate" and "Acetyl CoA" peaks, and a scale bar for 1000 units.

Table 5 (Cont'd)

from 1 to 666. Numbered from position 1.

Table 5 (Cont'd)

from 1 to 635. Numbered from position 1.

Figure 1 is a scatter plot with a fitted curve showing the relationship between the number of splits (S) on the x-axis and the number of genes (n) on the y-axis. The x-axis ranges from 0 to 10000 with major ticks every 1000 units. The y-axis ranges from 0 to 10000 with major ticks every 1000 units. The legend includes:

- Navy S = long ter (solid line)
- Navy S = short ter (dashed line)
- Navy S = 10000 (dotted line)

The data points show a non-linear relationship, starting at (0,0) and increasing rapidly before leveling off. A dashed line represents a linear fit to the initial data points. A dotted line represents the value n=10000 for all values of S.

Table 5 (Cont'd.)

from 1 to 6268. Numbered from position 1.

LAMINAR		1000+		2000+		3000+		4000+		5000+		6000+													
-Hairy 5' long ear																									
[Splits]																									
1 to 628 of 6268																									
Permeabilized																									
-Hairy 3' long ear																									
signal		1		1		1		1		1		1													
Scot221		1		1		1		1		1		1													
7612		1		1		1		1		1		1													
76031		1		1		1		1		1		1													
76032		1		1		1		1		1		1													
7611		1		1		1		1		1		1													
7603		1		1		1		1		1		1													
7602		1		1		1		1		1		1													
7601		1		1		1		1		1		1													
7600		1		1		1		1		1		1													
7602		1		1		1		1		1		1													
7601		1		1		1		1		1		1													
7600		1		1		1		1		1		1													
7602		1		1		1		1		1		1													
7601		1		1		1		1		1		1													
7600		1		1		1		1		1		1													
7602		1		1		1		1		1		1													
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7600		1		1		1		1		1		1													
7602		1		1		1		1		1		1													
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7602		1		1		1		1		1		1													
7601		1		1		1		1		1		1													
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7602		1		1		1		1		1		1													
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7602		1		1		1		1		1		1													
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7600		1		1		1		1		1		1													
7602		1		1		1		1		1		1													
7601		1		1		1		1		1		1													
7600		1		1		1		1	</																

Table 5 (Cont'd)

from 1 to 636. Numbered from position 3.

Table 5 (Cont'd)

from 1 to 655. Numbered from position 1.

from 1 to 6365. Numbered from position 1.

>NheI >Afl12
 mouse_DNA_end/HuMuSV_DNA_start_[Split]
 10 20 30 40 50 60 70
 TTTGAAAGAC CCCACCGTA GGTCGCAAGC TAGCTTAAGT AAAGCCACTT TCGAAAGCCAT GCAAAGATAC
 AAACCTTCTG GGGTCGGCAT CCACCGTTTG ATCGAAATTCA TTGGGGTGA AGCTTCGTTA CCTTTTATG

 >Pvu2
 >HpaB2 >EcoRS
 80 90 100 110 120 130 140
 ATAACTGAGA ATAGGAAGT TCACATCAG CTCAAGAACAA AAGAAACAGC TGAATACCAA ACAGGATATC
 TATGGACTCT TATGCTTCA AGCTCTAGTC CAGTOCTTGT TTCTTCTGCG ACTTATGGT TGTCTCTATA

 >Pvu2
 >HpaB2
 150 160 170 180 190 200 210
 TGTCGTAAGC GGTTCCTGCG CGGGCTCAAGC GCGAAAGAACAA GATGACACAG CTGACTGATG GCGCAACAG
 ACACCATTCG CCAAGGCGG CGCGCAGTCC CGGTTCTGT CTACTCTGTC GACTCACTAC CGGGTTTCG

 >ApaI
 >EcoRS >AluI >ApaI
 220 230 240 250 260 270 280
 GATATCTTG GTAGACAGTT CCTGGGGGGG CTGGGGGCGA AGAACAGATG GTCGGGAGAT GGGTTCTGAC
 CTATAGACAC CATTGCTCA GCACGGGGGG GAGGGGGGGT TCTTGTCTAC CAGGGGTCTA CGCCAGGTG

 >PpuXI
 >Pra2
 >Eco0109I
 >BamI >Styl >PstI >RsaI
 290 300 310 320 330 340 350
 CCTCACCAAGT TTCAAGTGAA TCATCACATG TTTCACAGGT GCGCCGAAAGG CCTGAAAGATG AGCCCTTATC
 GGAGTGTCA AGGATCACTT AGTACTCTAC AAAGCTCCCA CGGGGTTCTG GAGCTTCTAC TGGACATAGG

Table 6 (Cont'd)

>Ban2
 *
 >SacI
 *
 >SstI
 *
 >HpaII
 *
 360 370 380 390 400 410 420
 TATTTGAC TAACTAATCA GTTGGCTCTC CGCTTCGTTT CGCGGGCTTC CGCTCTCGGA GCTCAATATA
 ATAAACTTG ATTGGTATG CAGCGAAGA CGAAAGACAA CGCGCCGAG CGAGAGGGT CGAGTTATTC

 >Apa718
 *
 >Ban1
 *
 >SmaI
 *
 >AvaI
 *
 >Aql1 >RsaI
 *
 >Ban2 >BssH2 >HpaI >Tth111I >XbaI >RpaI
 430 440 450 460 470 480 490
 AGAGCCACCA ACCCTCTACT CGGGGGGCCA GTCTTCGGAT AGACTGGCTC CGCCGGGTAC CGTATTCGG
 TCTGGGGTGT TGGGGAGTGA CGGGGGCGGT CAGAGGCTA TCTGAGCGAG CGGGCCCGAT CGCATTAAGGG

 >StyI
 *
 500 510 520 530 540 550 560
 ATAAAGGCT CTTOCTGTTT SCATCGGAAT CGTGTCTCGT CTGTTCTTG CGAGGCTTC CTCTGAGTGA
 STATTCGGA GACCCACAA CGTGGCTTA SCACAGGACG GACAGGAAC CCTCCAGAG GAGACTCTACT

 >Ban2
 *
 570 580 590 600 610 620 630
 TTGACTACGCC AGCGGGGGCG TCTTCATTT CGGGGGTGT CGGGGATTTG GAGACCGCTC CGCGGGGCC
 AACTGATGGG TCCTGCCCCC AGAACTAAA CGCCCGACGA CGCGCTTAAC CTCTGGGAC CGCTCOCTGG

 >SalI

Table 6 (Cont'd)

Diagram of the restriction map of the 5' end of the 16S rRNA gene. The map shows restriction sites for Eco521, Cfr1, Xba3, Pst1, Sac1, and Sac1. The map is aligned with a sequence of 16S rRNA gene fragments from 780 to 840. The Sac1 site at position 810 is a double site.

>EcoNI
"
>Bsu36I
"
>AccI
"
>Sau3I
"
>Eco81I
"

Table 6 (Cont'd)

>Cvn1

.

>Met2

>Pst1

>Tth111I

850

860

870

880

890

900

910

CCCCACCTGA GCAAGGGAGT CGATGTCGAA TCCGACCCCG TCAGGATATG TGGTTCTGGT AGGRGACAG
GGCTGGACT CCTTCCCTCA GCTACACCTT AGGCTGGGGC AGTCCTATAC ACCAAGACCA TCCCTCTGTC

>Hpa1

920

930

940

950

960

970

980

AACCTAAAC AGITCCCGCC TCGCTCTGA TTTTTCGTTT CGCTTTCGAA CGAAGGCGC GCGCTCTGTC
TTGGATTTG TCAAGGGGG AGGCAGACTT AAAACGAAA CGCAAACTT GCGTTGGGG CGCGAACAG

>Pst1

990

1000

1010

1020

1030

1040

1050

TGCTGAGCA TGGTCTCTG TTGCTCTCT CTGACTGTGT TTCTGTATTT GTCTGAAAT TGGGCCAGA
ACGAGCTGGT ACCAGACAC AACAGACACA GACTGACACA AACACATAAA CAGACTTTA ATCCCGGTCT

>Koc1

.

>Sau1

.

>Cvn1

.

>Met2

.

>Ssu36I

.

>Afl2

>Eco81I

1060

1070

1080

1090

1100

1110

1120

CTGTTTACAC TCGCTTAAGT TTGACCTTAG GTCACTGGAA AGATGTCGAG CGGATGCGTC ACACCCAGTC
GACAATGGTG AGGAAATTCA AACTGGAAATC CAGTCACCTT TCTACAGCTC GCGTAGGGAG TGTGCGTCAG

>Cfr1

.

>AspAI

>Eas1

>Eas1

Table 6 (Cont'd)

>Mae2	>SstE2	>Pst1	>Bam1	>Kae2	>Cfr1
1130	1140	1150	1160	1170	1180
•	•	•	•	•	•
•	•	•	•	•	•
GGTAGATGTC	AAGAAGAGAC	CTTGGGTAC	CTTCCTCTCT	GCAGAATGCC	CAACCTTTAA
CCATCTACAG	TCTCTCTCTG	CAACCCATG	CAACAGAGA	CGCTCTTACCG	GGCCCTTACG
<hr/>					
>Bam1	>Mph1	>Pst1	>Bam1	>Kph1	>Cfr1
1200	1210	1220	1230	1240	1250
•	•	•	•	•	•
•	•	•	•	•	•
CGCGGAGAGC	GCACCTTTAA	CGAGACCTC	ATCACOCAGG	TTAAGATCAA	GCTCTTTCA
GGCGCTCTGC	CGTGGAAATT	GGCTCTGGAG	TAGTGGTCC	ATTCTGAGT	CCAGAAAACT
<hr/>					
>Pst1	>Dra2	>EcoO109I	>PpuMI	>Tch1112	>Sty1
•	•	•	•	•	•
•	•	•	•	•	•
ATGGACACCC	AGACCGAGTC	CCCTACATGG	TGACCTGGCA	AGCCCTGGCT	TTTGACCCCC
TACCTGCGG	TCTGGTCCAG	GGGATGTAGC	ACTGGACCTT	TOGGAACCGA	AAACTGGGGG
<hr/>					
>Rsa1	>Pst1	>Dra2	>EcoO109I	>PpuMI	>Tch1112
1340	1350	1360	1370	1380	1390
•	•	•	•	•	•
•	•	•	•	•	•
CAAGCCCTT	GTACACCCCA	AGCCCTGGCC	TCTCTCTCT	CCATCCCC	GCTCTCTCCC
GTTCGGAAA	CATGTGGGAT	TOGGAGGGG	AGGACAGAGCA	GGTAGGGGGG	GCAGAGAGGG
<hr/>					
>EcoN1	>EcoR1	>Pst1	>Dra2	>EcoO109I	>PpuMI
1410	1420	1430	1440	1450	1460
•	•	•	•	•	•
•	•	•	•	•	•
CTCTGTTGGA	CCCCCGCTG	ATCTCCCT	TATCGAGCC	TCACTCTCTC	TCTAAGGGGGG
GGACCAAGCT	GGGGGGGAGC	TACGACGGAA	ATAGGTTGGG	AGTCAGGAGA	AGATCCCC
<hr/>					

Table 6 (Cont'd)

>BspMI							>RsaI
1480	1490	1500	1510	1520	1530	1540	
•	•	•	•	•	•	•	
CTTGGTAAGT GACCAAGTAC AGTGGCAAAAC CATCAGCAAG CAGCTATGTA CTCTCCAGGG TGGGCGCTGGC							
GAACCAATTCA CTGGCTGATC TGACCTTTG CTAGCTGTTG CTCCATACAT GACAGCTTCC ACCGGGACGCG							
>RsaI							
1550	1560	1570	1580	1590	1600	1610	
•	•	•	•	•	•	•	
TTCCCCAGTC AACAGTCCAGG CGATTTGAGG GACGGCTTGGG GCTCTCTCT TACATGTACC TTTTGGTACG							
AAGGGGTACG TTCTGAGGTG CCTAAACTCC CTGGCACACCC CGAGAAGAGA ATGTACATGG AAACACGATCG							
>PstI							>HpaII
1620	1630	1640	1650	1660	1670	1680	
•	•	•	•	•	•	•	
CTCAACCCCTG ACTATCTTCC AGCTCATTTG TCCACATGG CCCTGTGCAAT CGACAGGTG CAACTCCGT							
GAATTTGGAC TGATAGAAGG TCCACTAACCA AGGTGTGACCC CGCACACCTA CCTGTCTTACG TTCTGAGACA							
>HpaI							
1690	1700	1710	1720	1730	1740	1750	
•	•	•	•	•	•	•	
CTTGCATTCG ACTAAGCTCT GCACCTTGCA CAACAGCTGC ACCTACTTCA AGTGTACAA AGAAACAGA							
GAACGTAACG TGATTCAGAA CGTGAACAGT GTTGTGACGG TGGATGAAGT TCAAGATGT TCTTTGCT							
>PvuII							
>HpaII							>HpaII
1760	1770	1780	1790	1800	1810	1820	
•	•	•	•	•	•	•	
GCTGCAACTG GACGATTAC TGTGGATTG ACACATGATT TTGAATGAA TTAAATGAA CGAAGATCC							
CGACGTTGAC CTGCTAAATG ACCACCTAA TGTCTACTAA AACTTACCTT AATTTAAATG TTCTCTTACGG							
>SphI							
>Hpa(7524)I							>PstI
1830	1840	1850	1860	1870	1880	1890	
•	•	•	•	•	•	•	
AAACTCAACCC CGATGGCTACG ATTTAAGTTT TACATGGCCA AGAAAGCCAC AGAAACTGAAA CGATGCGAGT							
TTTGAATGGG CGTACCGATG TAATTCGAA ATGTACACGGT TCTTCCCGTG TCTTGACTTT GTAGACGCTCA							

Table 6 (Cont'd)

>XbaI							>Afl12	
1900	1910	1920	1930	1940	1950	1960		
▼	•	•	•	•	•	•		
GTCTAGAAGA	AGAACTCAA	CCTCTGGAGG	AACTCTAAA	TTTACCTCAA	ACCAAAAACT	TTCACTTAA		
CAAGATCTCT	TCTTGAGTT	GGAGACCTCC	TTCACCGATT	AAATCCGATT	TTGTTTTTGA	AACTGAATTC		
							>AvaI	
							•	
>Ava2							>Aqui	
							•	
>SstI							>CcrI	
							•	
>SstI							>PacR7I	
							•	
>AatI							>Xba2	
							•	
1970	1980	1990	2000	2010	2020	2030	>BstY1	
▼	•	•	•	•	•	•		
GCCTAGGGAC	TTAATCAGCA	ATATCAAGCT	AAATGTC	CAGCTAAAGG	CATCTGAAAC	ACACATCAG		
CGGATCCCTG	AAATTAGTGT	TATAGTCA	TTATCAAGAC	CTGGATTTC	CTAGACTTTC	TCTTAAGTAC		
							>BstX1	
2040	2050	2060	2070	2080	2090	2100		
•	•	•	•	•	•	•		
TGTGAATATG	CTGATGAGAC	AGCCACATT	GTGGAATTTC	TGAACAGATG	GATTACCTTT	TGTCAUNGC		
ACACTTATAC	CACTACTCTG	TCGGTGCTAA	CAACCTTAAAG	ACTTGTCTAC	CTATCGAAA	ACAGTTTGT		
							>BamHI	
							•	
>BstY1								
							•	
>Xba2							>Hsp82	
							•	
>simian_virus_40_early_promoter								
							•	
>Mu-MuSV_DNA_end/simian_virus_40_DNA_start								
2110	2120	2130	2140	2150	2160	2170		
•	•	•	•	•	•	•		
TCATCTCAAC	ACTAACTTCA	TAATTAAGTC	CTTCCCACTT	AAACATATC	ACGATCCGCT	GTGGAATGTC		
AGTACAGTTG	TCATTGAACT	ATTAATTCAC	GAAGGGTCAA	TTTGTATAG	TTCTAAGGCGA	CACTTTAAC		
							>Bpu12II	
							•	
>HpaI								
							•	
>AvaJ								
							•	

Table 6 (Cont'd)

>Styl
-
>Avr2
-
>Stul
-
>BspMI

55

Table 6 (Cont'd)

>AatI	>KpnI	>PstI				
2460	2470	2480	2490	2500	2510	2520

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TCCAGAACTA GTGAGCCAGCC TTTTTGGAG CCCTAGGCTT TTGCAAAAGG CTTGGGCTGC AGGTGGAGGC
AGGTCCTCAT CACTCCCTCC AAAAACCTC CGGATCCCAA AACGTTTTTC GAACCCCAAGG TCCAGCTCCG

```

>BclI

>XbaI

>SstI

>BspMI

Imian_virus_DNA_end/Tn5_DNA_start

↓

↓

↓

↓

↓

↓

↓

↓

↓

↓

↓

```

2530 2540 2550 2560 2570 2580
↓ ↓ ↓ ↓ ↓ ↓
GGATCTGATC AACACACAGG ATCACCACTC TTTCGC ATG ATT GAA CAA GAT GGA TTG CAC GCA GCT TCT
GCTAGACTAG TTCTCTCTCC TACTCTAGC AAAGCG TAG TAA CTT GTT CTA CCT AAC GTC GGT CCA AGA
Met Ile Glu Gln Asp Gly Leu His Ala Gly Ser>

```

>Eco52I

>EagI

>Esel

>CfrI

>XbaI

Table 6 (Cont'd)

>Ban1		>Pst1		>Eco1		>Cfr1		>Sal1											
2720	2730	2740	2750	2760	2770	2780	2790	2800	2810										
TCG	GGT	GCC	CTG	AAT	GAA	CTG	CAG	CAC	CAO	GCA	GCG	CGG	CTA	TGG	CTG	GCC	ACG	ACG	GGC
AGG	CCA	CCG	GAC	TTA	CTT	GAC	CTC	CTG	CTC	OCT	CCG	GAT	ACC	ACC	CGC	GGG	TGC	TCC	CCC
Ser	Gly	Ala	Leu	Asn	Cys	Glu	Cys	Asp	Glu	Ala	Ala	Arg	Leu	Ser	Trp	Leu	Ala	Thr	Gly

```

>Fsp1
-
>Acl1
-
>Tth111I
-
>PvuII
-
>Pfd12
-
>HglA1
-
>MspI
-
>NspI2
-
>HaeII
-
2780  | 2790 | 2800 | 2810 2820 2830 2840
*   *   *   *   *   *   *
GTT CCT TCG GCA CCT CTC CTC GAC GTC ACT GAA GCG GCA AGG GAC TCG CTA TTG GGC
CAA CGA AGG CGT GCA CAC CGC CTC CAA CAG TCA CTT CGC CCT TCC TCG ACT GAC GAG GAT AAC CGC
Val Pro Leu Ala Val Leu Asp Val Val Thr Cys Ala Gly Arg Leu Ser Thr Leu Leu Gly

```

```

>BstYI
.
.
>Ehol >Mph1
.
.
2850 | 2860 | 2870 2880 2890 2900
.
.
CAA GTC CGG GGG CAA GAT CTC CTC TCA TCT CAC CTT CCT GCT CCT GGC GAG AAA GTC TCC ATC ATG
CTT CAC CGG CCC GTC CTA GAG GCA AGT AGA GTC GAA CGA CGG CGG CGG CTC TTZ CAZ AGG TAG TAC
Glu Val Pro Glu Glu Leu Leu Ser Ser His Leu Ala Pro Ala Glu Leu Val Ser Ile Met

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Table 6 (Cont'd)

2910	2920	2930	2940	2950	2960
GCT GAT GCA ATG CGC CGG CTC CAT ACG CTT GAT CGG GCT ACC TCC CCA TTC CAC GAC CAA CGG					
CGA CTA CGT TAC CGC CGC GAC CTA TGC CAA CTA CGC CGA TGG AGG GGT AAG CTG CTG GTT CGC					
Ala Asp Ala Met Arg Arg Leu His Thr Leu Asp Pro Ala Thr Cys Pro Phe Asp His Cln Ala>					

>Rsa1

>MgIAl

>Kpn2

>Cfr10I

2970	2980	2990	3000	3010	3020	3030
AAA CAT CGC ATC GAG CGA GCA CGT ACT CGG ATG GAA CGC CGT CTT GTC GAT CAG GAT GAT CTC						
TTC GTA CGG TAG CTC CGT CGT CGA TGA CGC TAC CTT CGG CGA CGA CAG CTA CTG CTA CTA GAC						
Lys His Arg Ile Glu Arg Ala Arg Thr Arg Met Glu Ala Glu Leu Val Asp Cln Asp Asp Leu>						

>SphI

>Msp(7524)1

>Bam2

>BssH2 >MspI

3040	3050	3060	3070	3080	3090
GAC GAA CAC CAT CAC CGC CTC CGC CGA CGC GAA CTC TTC CGC ACG ACG CTC AAC CGG CGC ATG CGG					
CTG CTT CTC CTA GTC CGC CGC CGT CGG CTT GAC AAG CGG TCC GAG TTC CGC CGC CGC TAC CGG					
Asp Glu Glu His Cln Glu Leu Ala Pro Ala Glu Leu Phe Ala Arg Leu Lys Ala Arg Met Pro>					

>Xba2

>Nco1

>Cfr1

>BstY1

>Sty1

>Eae1

3100	3110	3120	3130	3140	3150
GAC CGC GAG GAT CTC GTC GTC ACC CAT CGC GAT CGC TGC TTC CGG AAT ATC ATG GTC GAA AAT					
CTG CGG CTC CTA GAG CGG CGC CTA CGG CGC AAC AAC CGC TTA TAG TAC GAC CGT TTA					
Asp Gly Glu Asp Leu Val Val Thr His Gly Asp Ala Cys Leu Pro Asn Ile Met Val Glu Asn>					

Table 6 (Cont'd)

>Cfr10I

>Cfr1

>Eco1

>Hae1

>Rsr2

3160	3170	3180	3190	3200	3210
.
GGC CGC TTT TCT GGA TTC ATC GAC TGT	GGC CGG CTG GGT	GTG CGG GAC GGC TAT	GAG GAC ATA		
CGG CGC AAA AGA CCT AAG TAG CTG	ACA CGG GCG GAC CCA	CAC CGC CTG CGG ATA	ATC CTC TGT TAT		
Gly Arg Phe Ser Gly Phe Ile Asp Cys	Gly Arg Leu Gly Val	Ala Asp Arg Tyr	Gly Asp Ile		

3220	3230	3240	3250	3260	3270	3280
.
CGG TTG GCT ACC CGT GAT ATT	GCT GAA GAG CTT CGC	GGC GAA TGG GCT	GAC CGG TTC	CTC CTC		
CGC AAC CGA TGG GCA CTA TAA CGA	CTT CTC GAA CGG CGG CTT ACC CGA CTG CGG AAG	CAC GAC				
Ala Leu Ala Thr Arg Asp Ile	Ala Glu Glu Leu Cys Glu Trp	Ala Asp Arg	Phe Leu Val			

3290	3300	3310	3320	3330	3340
.
CTT TAC GGT ATC GGC GCT CCC CAT	TOC CAG CGC ATC GGC TTC TAT	GGC CTT CTT GAC GAG TGC			
GAA ATG CCA TAG CGG CGA CGG CTA AGC GTC CGG TAG CGG AAG ATA CGG GAA GAA CTC CTC AAG					
Leu Tyr Gly Ile Ala Ala Pro Asp Ser Cys Arg Ile	Ala Phe Tyr Arg	Leu Leu Asp	Gly Phe		

>Pst1

>Tns_DNA_end/_No-MuLV_DNA_start

3350	3360	3370	3380	3390	3400	3410	3420
.
TTC TGA CGGGGACTC TCGGGTTGCA	TAATATATAA CATTITATTG	AGCTCTCCAGA	AAAAGGGGGG	AATGAAAGAC			
AAG ACT CGCCCTGAG ACCCCGAGCT	ATTTTATTTT	CTAAATATAA	TCAGGGCT	TTTTCGGGGG	TTACTTTCTG		
Phe End>							

>Afl12

>Whe1

3430	3440	3450	3460	3470	3480	3490
.
CCGACCTGTA CGTTTGGCAA CGTACGTTAA	GTAAACCCAT	TTTCCGAGG	ATGGAAAAAT	ACATAACTG		
GGCTGGACAT CCAGGCGCTT CGATCGAAAT	CAATGGGCGTA	AAACGTTGCG	TACCTTTCGA	TGTATGACT		

>Nsp82

>Pvu2

>EcoRI5

3500	3510	3520	3530	3540	3550	3560
.
GAATAGAGAA GTTCAGATCA	AGGTCAGGAA	CACATGAAAC	ACCTGAAATAT	GGGCGAAACA	CGATATCTGT	
CTTATCTCTT	CAAGCTGACT	TCAGCTGCTT	GTCTGACCTT	CCCGGTGTTG	CTTATAGAGAC	

>AlwN1
 3640 3650 3660 3670 3680 3690 3700
 TCTCTGGTAA GCGAGTTCCTC CCGCGCTCA GGGCCAAAGCA CAGATGGTCC CCAGATGGCG TCCACCCCTC
 AGACACCAATT CCTCTACGGC GGGGGCGAGT CCGGGCTCTC GCTCTACCCAGG GGTCTAACCC AGGTCCCGAC

>Sac2	>Ban1
.	.
>Bba1	>Acp718
.	.
>Ple1	>Ple1
.	.

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Table 6 (Cont'd)

>Mar1 | >Sma1 |
 * | * |
 * | * |
 >Bam1 | >Xba1 |
 * | * |
 * | * |
 >Apa1 | >Acy1 | >Ava1 | >Kpn1
 * | * | * |
 * | * | * |
 >Bam2 | >Ava1 | >Aha2 | >Tct1111 | >Apa1 | >Rsa1
 | 3850 | 3860 | 3870 | 3880 | 3890 | 3900 | 3910
 | v | v | v | v | v | v | v
 CCACACACCC TCACTCGGG CGCCAGCTCT COGATTGACT GAGTGGCCCG GGTAACCGTG TATCCAAATA
 GGTGTTGGGG AGTCAGCCCC CGCGTCAGGA GGCTTAACGTG CTACGGGGC CCATGGGCAAC ATAGGGTTATT

 >Sty1
 3920 3930 3940 3950 3960 3970 3980
 * * * * * * *
 ACCCTCTTGC AGTTGCATC GACTTGTGGT CTGGCTGTTC CTGGGAGGG TCTCTCTGA GTGATTCGAC
 TCGGAGAAC TCAAGCTAGG CTGAAACACCA GAGGCCAACG GAACTCTCC AGAGGAGACT CACTAACTCA

 >Nsp82 >Bam2
 3990 4000 4010 | 4020 4030 4040 4050
 * * * * * * *
 ACCCGTCAGC GGGGGCTTT CATTGGGGG CTGGCTGGGG ATGGGAGAC CCCTGGCCAG GGACCCACCA
 TCGGCACTGG CGGGCAAGAA GTAAACCCCC GAGCAGGGGG TAGCCCTCTG GGGACGGCTC CCTGGTGGGT

 >Nsp1752
 >Rph1 >Rph1 >Rph1
 4060 4070 | 4080 4090 4100 | 4110 4120
 * * * * * * *
 >Mo-MuLV_DNA_end/plasmid_pBR322_DNA_start
 4060 4070 | 4080 4090 4100 | 4110 4120
 * * * * * * *
 CGGACCCACCG CGGCTTAACG TGGCTGGCTC CGGGCTTTCG GTGATGACCG TGAACACAGC TGACACATCG
 GGGTGGTGGC CCTCTATGG ACCGAGGGAG CGGGCAAGAC CACTACTGC ACTTTGGAC ACTTGTACG

 >Rpa1
 4130 4140 4150 4160 4170 | 4180 4190
 * * * * * * *
 AGCTCCCCCA GAGGCTTACG CTGTCCTGT AAGGGGATGC CGGGAGCAGA CAGGGGCTC AGGGGGCTC
 TGGAGGGCTC CTGGCTGTG CGAACACAGA TTGGCTTACG GGGCTGGTCT GTGGGGGCGA TGGGGGGCGA

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Table 6 (Cont'd)

	4200	4210	4220	4230	4240	4250	4260
ACGGGGTGTGTT	GGGGGGTGTGTC	GGGGCCCAAC	CATGACCCAG	TCACGGTAGCG	ATACGGCAGT	GTATACGTGCG	
TOGCCACAA	CCGGCCACAG	CCCCGGCTCG	GTACTGGTC	ACTGCATGCC	TATGGCTCA	CATATGACCC	

>Hg1A1

	>Real			>ApaI	>NdeI		
	4270	4280	4290	4300	4310	4320	4330
TTAACTATCG	GGCATCAGG	CAGATTTGAC	TCACAGTGC	CCATATGGCG	TCTGAAATAC	CCACACAGTG	
AATTGATAAC	CGCTGACTTC	GTCTAACATG	ACTCTCACTG	GTATATGCC	ACACTTTATG	GGGTGTCAC	

>HaeII

>PstI

	4340	4350	4360	4370	4380	4390	4400
CGTAAGGAGA	AAATACCGCA	TCAGGGGCTC	TTCCGCTTGC	TCGCTCACTG	ACTGGCTGG	CTGGGTGTT	
GGATTCCTCT	TTTATGGGT	AGTGGGGAG	AGGGCGAAGG	ACCCGAGTGAC	TGAGGGAGCC	GACCCACCAA	
4410	4420	4430	4440	4450	4460	4470	
CGGCTGGGC	GACGGGTATC	AGCTCACTCA	AAAGGGTAA	TACGGTTATC	CACAGAAATCA	GGGGATAACG	
GGCCACCCGCG	CTCCUCCATAG	TCGAGTCAGT	TTCCGGCATT	ATGCCAATAG	CTGTCTTACTG	GGGGATTCG	

>Hsp(7824)1

>HspM1

>Afl13

	4480	4490	4500	4510	4520	4530	4540
CAGGAAGAA	CATGTGACCA	AAAGGCCAGC	AAAGACCCAG	GAACCGTAA	AAAGGGGGGT	TCTGGGGTT	
GTCTTCTT	GTACACTGT	TTTGGGTG	TTTTCGGTC	CTTGGCATTT	TTCCGGGCGA	ACGACCGAA	

>HpaI

	4550	4560	4570	4580	4590	4600	4610
--	------	------	------	------	------	------	------

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Table 6 (Cont'd)

TTTCCATAGG CTCGGGGGGG CTGACCCAGCA TCAACAAAAAT CGACGCTCAA GTGACAGCTG CGCGAAACGGG
AAAGGTATGC GAGGGGGGGG GACTGCTGCT AGCTGTTTTA GCTGCGACTT CAGTCCTCAC CGCTTTGGCC

4620 4630 4640 4650 4660 4670 4680

ACAGGACTAT AAACATACCA CGCGTTTCCC CCTCGGAAGCT CGCTGTGCG CTCTCTGTT CGGACACCTGC
TGTCTCTGATA TTTCATGCT CGCCAAAGGC CGACCTTGA CGGACCAACGC GAGAGGAAAC CGCTGGGACG

>Hae2

4690 4700 4710 4720 4730 4740 4750

CGCTTACGGG ATACCTGTCG CGCTTCTCC CTTCGGGAGG CGTGGCGCTT TCTCATAGCT CACGCTTGTG
GCCAAATGGCC TATGGACAGG CGGAAACAGG GAAGCCCTTC GCACCGGGAA AGACTATOGA GTGCGACATC

>HgIAl

>ApalI

4760 4770 4780 4790 4800 4810 4820

GTAATCTCACT TCCGCTGAGG TGGTTCGCTC CAAGCTGGGC TGCTGTGCAAC AGCCCCCGT TCAGCCCCGC
CATAGAGTC AAGCCACATCC ACCAGGGAG GTTGGACCCCG AGCACACGTG TGCCCCGGCA AGTGGGGGCTG

>Nsp82

>PstI

4830 4840 4850 4860 4870 4880 4890

CGCTTGGCGT TATCCGGAA CTATGCTCTT GAGTCCAAACC CGCTAACACA CGACTTATGCG CGACTCGCG
GGAGCCGGCA ATAGGCCATT GATAGGCAAA CTCAAGGTTGG CGCATTCTGT GCTCAATAGC CGTGACCGTC

>AluNI

4900 4910 4920 4930 4940 4950 4960

CAGCCCACTGG TAACAGGATT ACCACAGCGCA CGTATGCTAGG CGCTGCTACA CAGTCTCTGA AGTGGTGGGC
GTCGGTGACCC ATTGCTCAA TGGTCTGCT CGCATACATCC GCCACGATGT CTCAAGAACT TCAACACCGG

4970 4980 4990 5000 5010 5020 5030

TAACCTACGGC TACACTAGAA CGACAGTATG TGCTATCTGC CCTCTGGCTGA ACCGACTTAC CTTCGGGAAA
ATTGATGCCG ATCTGATCTT CCTCTCTCAA ACCATAGACGG CGAGACGACT TGGTCAATG GAAGCCCTTG

>Nsp82

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Table 6 (Cont'd)

PCT/US92/08999

5040	5050	5060	5070	5080	5090	5100
•	•	•	•	•	•	•
ACAGTTGGTA	GCTCTTGTAC	CCGCAACCAA	ACCACCGCTG	GTAGGGCTGG	TTTTTTGTG	TGCAACGAC
TCTCAACCAT	CGAGAACATG	CCCGTTTGT	TGCTGGGAC	CATCCCCACC	AAAAAAACAA	ACGGTGGCTG

>Xba2 >BstYI

5110	5120	5130	5140	5150	5160	5170
•	•	•	•	•	•	•
>BstYI	>Xba2					>RgaI

AGATTAACGG	CAGAAAAAAA	GGATCTCAAG	AGAGATCTT	GATCTTTCT	ACGGGGCTCG	ACCGTCAGTC
TCTAAATGCGC	GTCTTTTTT	CTCTAGAGTC	TCTCTAGAAA	CTAGAAAAAGA	TGCCCCAGAC	TGGCAGTCAC

>BstYI

5180	5190	5200	5210	5220	5230	5240
•	•	•	•	•	•	•
>Mae2	>BspMI		>HpaI	>Xba2	>BstYI	>DraI

GAACCAAAAC	TCACTTAAAG	GGATTTGGT	CATAGATTA	TCAAAAAGGA	TCTTCACCTA	GTCCCTTTTA
CTTGGCTTTG	AGTGCATTTC	CCTAAACCA	GTACTCTAAT	AGTTTTCCCT	AGAAGTGGAT	CTAGGAAAT

>DraI

>Aba3

5250	5260	5270	5280	5290	5300	5310
•	•	•	•	•	•	•
ATTTAAAAAT	CAAGTTTTAA	ATCAATCTAA	ACTATATATG	AGTAAACTTG	GTCTGACAGT	TACCAATGCT
TTAATTTTTA	CTTCAAATT	TAGTTAGATT	TCATATATAC	TCATTTGAAAC	CAGACTGTCA	ATGGTTTACGA

>BsrI

>PstI

5320	5330	5340	5350	5360	5370	5380
•	•	•	•	•	•	•
TAATCAGTCA	GGCACTTATC	TCAGGAGTC	GTCTTATTTG	TCATCCATA	GTGGCTGAC	TCGGGGTGTG
ATTAGTCACT	CGCTGATAG	AGTGGCTAGA	CAGATAACG	AACTAGGTAT	CAACGGACTG	ACGGGGCGCA

>BpuI

5390	5400	5410	5420	5430	5440	5450
•	•	•	•	•	•	•
GTACATAACT	ACGATAACGG	ACGGCTTACG	ATCTGGGGCG	AGTCTCTGAA	TGATACGGCG	ACGACGAGCG
CATCTATGCA	TCCTATGCC	TCGGGATGG	TAGACGGGG	TCAGGACGTT	ACTATGGCGC	TCTGGGTGCG

>Cfr10I

>BglII

5460	5470	5480	5490	5500	5510	5520
•	•	•	•	•	•	•
TCACCGGCTC	CAGATTATTC	ACCAATAAC	CAGCCGACCG	GAAGGGCCGA	GGCCAGAAGT	GTGCTTGCAA
AGTGGGGCG	GTCTAAATAG	TGTTTATTC	GTGGCTGGC	CTTCCCCGGT	CGCGTCTTC	CGGGCGGTT

Table 6 (Cont'd)

>HaeI

5530	5540	5550	5560	5570	5580	5590
CTTTATCCGC	CTCCATOCAG	TCTATTAATT	GGTGGCCCGA	ACCTAGAGTA	AGTAGTTCGG	CAGTTTAATG
GAATAAOGCG	GAGGTAGGTC	AGATAATTAA	CAAOGCCCT	TGATCTCAT	TGATCAAGCG	GTCATAATTG

>MaeI

>AosI

>PspI

>FdiI2

>HstI

>PstI

5600	5610	5620	5630	5640	5650	5660
TTTGGCCGAAAC	TTTGTGCGCA	TTGCTGCCAGG	CATOGTGTG	TCAOGCTGT	CGTTTGGTAT	GCCTTCATTC
AAACGCCCTTG	CAACAAOGGT	AAAGCAOGTCC	GTACCCACAC	AGTGGCCAGCA	GCACACCCATA	CGGAGTGTAG

5670 5680 5690 5700 5710 5720 5730

AGCTCCGGTT	CCCAAGATC	AGGGCGAGTT	ACATGATCC	CCATGTTGTG	CAAAAGCG	GTAGCTGT
TOGAGGGCCAA	GGGTGCTAG	TCCCGCTAA	TGACTAGGG	GGTACACAC	GTTCCTTCCC	CAATCGAGGA

>PvuI >EaeI

>KorI >CfrI

5740	5750	5760	5770	5780	5790	5800
TCGGCTCTCC	CATCGTTTGC	AGAAGTAAGT	GGCCCGACT	GTATACACTC	ATGGTTATAG	CAGGACTCGA
AGCCAGGAGG	CTTACCAAG	TCTTCATCA	ACCGGGCTCA	CAATAGTGAG	TACCAATAACG	GTGTTGAGGT

>RsaI

>ScaI >RphI

5810	5820	5830	5840	5850	5860	5870
TAATTCCTCT	ACTGTCATCG	CATCCGTAAG	ATGCTTCTTG	GTGACTGGTG	AGTACTCANC	CAACTCAATC
ATTAACAGAA	TGACAGTAGG	GTAGGCGATTC	TAAGAAGAGA	CACTGACAC	TCATGAGTTG	GTTCAGTGAAG

Table 6 (Cont'd)

GAATACTCAT ACTCTTCTT TTCAATATT ATTCAAGCAT TTATCAGGT TATTTCTCA TGACCGGATA
 TTATCAGTA TGACAGGAA AAGCTTATAA TAATCTGTA AATAGCTCCA ATAACAGAT ACTCGCTTA

6230 6240 6250 6260 6270 6280 6290
 CATATTTGAA TGATTTTAA AATAATTAACA AATAGCGGTT CGGGCCACAT TTCCCCCAAAG AGTGCACCT
 GTATAAACTT ACATAAACTT TTATTTTGT TTATCCCAA GGCGCGTGTAA AGGGGGCTT TGACGGTGGAA

>Aat2
 >Aha2
 >Acy1
 >Hae2
 >BspRI
 >Pst1
 >EcoO109I
 >Dra2
 6300 6310 6320 6330 6340 6350 6360
 CACCTCTAAG AACCATTAT TATCATGACA TTAACTTATA AAAATAGGCG TATCAGGAGG CGCTTTCGTC
 CTGGAGATTC TTGGTAAATA ATAGTACTGT AATGGATAT TTATCCCG ATAGTGTTC CGGAAAGGAG

TTCAA
 AAGTT

zymes which do not cut LXSNRIZL2 :

Hcc3	Bgl2	Cla1	Npa1	Nru1	SnaB1
Hpa1	Bsm1	Dra3	Mlu1	PflM1	Sph1
Hpa2	BspM2	Eco47III	Mre1	Sac2	Sst2
Hpa3	BstS1	Esp1	Not1	Sall	

To generate the LXSN-RI-IL2 retroviral vector, 10 micrograms of pLXSN-RI-IL2 DNA was transfected into the ecotropic packaging cell line PE501 by standard calcium phosphate precipitation methods (Miller et al., Mol. Cell Biol. 6:2895, 1986). The transfected PE501 cell line was grown in DMEM medium with 10% FCS. The medium was changed after 24 hours and supernatant harvested 24 hours later to infect the amphotropic packaging cell line PA317, as described (Miller et al., Mol. Cell Biol. 6:2895, 1986 and 10 Miller et al., BioTechniques 7:980, 1989). The infected PA317 cells were harvested by trypsinization 24 hours later and replated 1:20 in DMEM containing 10% FCS and the neomycin analogue G418 (400 µg/ml). The cells were grown at 37°C in 7% CO₂ atmosphere. The selection medium was 15 changed every 5 days until colonies appeared. On day 14, twenty colonies were selected, expanded and tested for viral production by standard methods (Xu et al., Virology 171:331-341, 1989). Briefly, supernatants were harvested from confluent culture dishes, passed through a .45 µm 20 filter, diluted with DMEM with 10% FCS and utilized to infect NIH 3T3 cells in the presence of 8 µg/ml polybrene. After 24 hours, the infected NIH 3T3 cells were grown in culture medium that contained the neomycin analogue G418. After 12-14 days, the colonies were stained, counted and 25 the viral titer calculated as described (Xu et al., Virology 171:331-341, 1989).

Colonies with the highest viral titers (>10⁴ infectious units/ml) were tested for IL-2 expression by Northern blot analyses. Colonies with the highest viral 30 titers and documented IL-2 expression were cryopreserved and will be utilized as stock cultures to produce the LXSN-RI-IL2 retroviral vector trial.

EXAMPLE IVRETROVIRAL VECTOR CONSTRUCTION AND CYTOKINE EXPRESSION

To increase IL-2 production by transduced cell lines, vectors were used containing different promoters to 5 drive IL-2 expression, and a human IL-2 cDNA was directionally sub-cloned into the insulin secretory signal peptide (17). The IL-2 cDNA was directionally sub-cloned into the parental plasmids of the LXSN (LTR promoter) and LNCX (CMV promoter) vectors (gifts of Dr. A.D. Miller) 10 (18). The newly constructed vectors (Figure 1), designated as LXSN-IL2 and LNCX-IL2, were packaged in the PA317 cell line for production of retroviral supernatant. As a control, the high level expressing, double copy vector DC/TKIL-2 vector (thymidine kinase promoter) (a gift of Dr. 15 E. Gilboa) was used for comparison.

These vectors were used to transduce a number of murine and human, primary and established cell lines. Pools of transduced cells were selected and expanded in DMEM medium, containing 10% fetal bovine serum (FBS) and 20 400 µg/ml of active G-418, a neomycin analogue. The results of expression studies in the MCR9 and Balb/c 3T3 cell lines are presented in Table 7.

Table 7

Comparison of IL-2 expression by fibroblasts
transduced with different IL-2 vectors.

5

Fibroblast	Vector	<u>ng IL-2</u>	<u>Units IL-2</u>
		per 10 ⁴ cells per day	
Murine	LNCX (Control)	0.4 ±50%	<1
	LNCX-IL2	33.7 ±11%	67
	LXSN-IL2	6.6 ± 6%	13
	DC/TKIL-2	1.9 ± 5%	4
Human	LXSN (Control)	0.7 ±29%	1
	LNCX-IL2	159.5 ±17%	319
	LXSN-IL2	25.5 ±15%	51
	DC/TKIL-2	3.0 ±10%	6

EXAMPLE VFIBROBLAST CULTURE AND CONDITIONS FOR RETROVIRAL
TRANSDUCTION

The culture conditions for the growth of primary
5 fibroblasts retroviral transduction were optimized.
Primary fibroblasts were successfully cultured. The
optimal conditions enable the growth of approximately 3-4
x 10⁶ primary fibroblasts from a 12 mm² skin biopsy in
approximately 4-6 weeks. Retroviral infection, G418
10 selection, and expansion of the genetically modified
fibroblasts takes an additional 4-6 weeks.

Exploring the conditions for genetic modification
of primary fibroblasts suggests that optimal transduction
may be obtained by the following procedure: The fibroblasts
15 are synchronized in G1 phase by serum starvation, followed
by stimulation with medium containing 15% fetal bovine
serum 15 hours prior to transduction. The cells are then
subjected to 2 cycles of retrovirus infection, each cycle
lasting approximately 3 hours. The cells are refed with
20 fresh media overnight, and then selection in G418 is
initiated the next day. This method is capable of
transducing 5-15% of the fibroblasts in a culture,
depending on the multiplicity of infection.

This procedure was used to transduce a large
25 number of primary and established fibroblasts. As an
example, Table 8 compares the expression levels of IL-2 in
fibroblast lines transduced with LXSN-IL2.

Table 8

Expression of IL-2 by fibroblasts transduced with LXSN-IL2.				
5	Fibroblast		ng IL-2 Units IL-2	
	Line	Species	Origin	per 10 ⁶ cells per day
10	Balb/c 3T3	Murine	Transformed	6.6 ± 6% 13
	MCR9	Human	Embryonic	25.5 ± 15% 51
	NHDF 313	Human	Skin	25.0 ± 10% 50
	GT1	Human	Skin	15.0 ± 5% 30

These results indicate that the IL-2 expression levels in established, embryonic, and primary fibroblast cultures are similar. Comparison of these data with Table 7 suggest that IL-2 expression is affected more by factors such as different promoters than by the fibroblast line used. Similarly, changes in culture conditions can have important effects on IL-2 expression. Table 9 shows that transduced GT1 cells, a primary human fibroblast culture expressed 15-fold more IL-2 under 100 µg/ml G418 selection than under 25 µg/ml G418 selection. Several other primary fibroblast lines have also been transduced with our vectors and are currently growing under G418 selection.

Table 9

Effect of G418 concentration on IL-2 expression by GT1 cells transduced with LXSIL2.		
5	Selection dose of G418	ng IL-2 secreted per 10 ⁶ cells per day*
10	25 µg/ml	1.0 ± 10%
	50 µg/ml	3.0 ± 6%
	100 µg/ml	15.0 ± 5%

*After three weeks of G418 selection.

EXAMPLE VI

15 COMPARISON OF IL-2 EXPRESSION LEVELS INDUCED
PERIPHERAL BLOOD LYMPHOCYTES AND
GENETICALLY MODIFIED FIBROBLASTS

In order to compare the production of IL-2 by genetically modified fibroblasts to that achieved by 20 stimulating normal human peripheral blood lymphocytes (nPBL) in vitro, nPBL were isolated by Ficoll-Paque density centrifugation, and cultured in the presence of allogeneic nPBL (mixed lymphocyte culture, MLC) or 2 µM calcium ionophore (CI) (A23187) free acid) plus 17 nM phorbol 12-25 myristate 13-acetate (PMA). The results of this experiment, present in Table 10, indicate that the level of IL-2 expression in the PMA/CI stimulated normal T cell population was 2 ng/10⁶ cells/24 hours. This is equivalent to IL-2 expression by Balb/c 3T3 fibroblasts transduced 30 with DC/TKIL-2 (Table 7), our least productive vector. The level of IL-2 expression in the MLC was 130 pg/10⁶ cells/24 hours. This was lower than the PMA/CI stimulated culture, presumably because PMA/CI induced a nonspecific response

while MLC resulted in specific Th stimulation. When the estimated percentage of antigen-specific Th in the MLC-stimulated population is taken into consideration, the level of IL-2 expression per stimulated T cell becomes 5 equivalent for both methods.

Table 10
Levels of IL-2 secretion by different cells.

	Cells	pg IL-2 secreted per 10 ⁶ cells per day
10		
	Lymphocytes:	
	Control (non-activated)	5 ± 50%
	PMA + Calcium Ionophore	2,000 ± 6%
15	Mixed lymphocyte culture	130 ± 90%
	Transduced fibroblasts:	
	MCR9-LXSN-IL2	24,000 ± 5%
	MCR9-LNCK-IL2	162,000 ± 20%
	MCR9-DC/TKIL-2	10,000 ± 6%
20		

EXAMPLE VII

FIBROBLAST MEDIATED CYTOKINE GENE THERAPY
IN MURINE TUMOR MODELS

Two experimental protocols were used to study the 25 efficacy of fibroblast-mediated cytokine gene therapy on induction of anti-tumor immunity. The first protocol was designed to test the effects of genetically modified fibroblasts on tumor implantation, while the second protocol was designed to induce a systemic anti-tumor 30 immunity. The results of each experiment are presented with two figures and one table. In the first figure, the rate of tumor growth for each treatment group is presented

as the mean tumor size in the group over time. In the second figure, a Kaplan-Meier curve presents the time of tumor onset for the individual animals in each treatment group. The number of animals, the number and percentage of 5 tumor free animals, and the tumor size distribution patterns for each experiment are presented in a table.

EXAMPLE VII(a)

EFFECT OF FIBROBLAST MEDIATED CYTOKINE GENE
THERAPY ON TUMOR IMPLANTATION

10 Mice were injected subcutaneously with mixtures of 5×10^4 CT26 cells and 2×10^4 fibroblasts genetically modified by different retroviral vectors to express IL-2. In the control arms injected with tumor cells only, or with tumor cells mixed with unmodified fibroblasts, 31 of 33 15 animals (94%) developed tumors by 4 weeks (Figures 6 and 7, Table 9). In contrast, 22 out of the 34 animals (65%) receiving fibroblast mediated cytokine gene therapy were tumor free at 3 weeks, and 5 animals (18%) remain tumor free after 12 weeks. Those animals that received 20 fibroblast mediated IL-2 therapy and developed tumor were characterized by a delayed onset and rate of tumor growth.

Table 11

Effect of IL-2 modified fibroblasts on tumor establishment and development.
 2×10^6 fibroblasts mixed with 5×10^4 CT26 tumor cells at time of injection.

Fibroblasts mixed with tumor cells	Animal Number	Tumor- bearing	Percent Tumor-free	Tumor Size (mm ²)			Median Tumor Size (mm ²)
				25-100	101-200	>301	
<i>After 12 Weeks*</i>							
Control (no fibroblasts)	11	0	11	0%	1	0	9
Unmodified fibroblasts**	13	2	11	15%	1	0	7
DCTK-IL2 fibroblasts	13	0	13	0%	1	3	4
LNCK-IL2 fibroblasts	13	5	8	39%	5	2	0

* Mean tumor size is for 4 weeks, the last timepoint at which tumors were measured.

** Two mice in this arm developed intraperitoneal tumors which were not measurable.

After 3 weeks the mean tumor size (measured as the product of the longest and widest tumor axes) in the control group of mice was 128 mm², compared to 68 and 7 mm² in groups of mice injected with tumor cells mixed with 5 fibroblasts transduced with DC/TKIL-2 or LNCX-IL2, respectively. This resulted in a highly significant difference (corrected $\chi^2 = 18.69$, $p = 0.001$) between the IL-2 treated animals compared to the mice treated with CT26 alone or CT26 mixed with unmodified fibroblasts. After 10 four weeks the equivalent measurements were 373,300 and 72 mm² (Table 11). It is notable that LNCX-IL2, the highest expressing vector caused substantially greater inhibition of tumorigenicity than the lower expressing vector DC/TKIL-2. A multivariate non-parametric statistical procedure 15 (19,20), utilized to evaluate differences in tumor growth, demonstrated that after 4 weeks the differences between the growth curves for the four groups presented in Figure 2 were highly significant ($p < 0.001$). Subsequent comparisons between the control arm and animals that 20 received tumor cells mixed with IL-2 transduced fibroblasts revealed a significant difference ($P < 0.05$). The differences between the animals injected with tumor cells alone, and those injected with tumor cells plus unmodified fibroblasts were not significant, while the differences 25 between animals receiving low IL-2 expressing fibroblast, and those receiving high IL-2 expressing fibroblasts was significant ($P = 0.05$).

When mice were injected with 2×10^6 modified 30 fibroblasts mixed with 1×10^6 live tumor cells the results became more striking (see Figures 8 and 9, and Table 12). All the control animals developed tumors after 4 weeks whereas 33% and 27% of the animals treated with fibroblasts modified with the DCTK-IL2 or LXSN-IL2 vectors (respectively) remain tumor free after 7 weeks (the 35 experiment is ongoing). More dramatically, 75% of the animals treated with fibroblasts modified with the highest

IL-2 producing vector, LN CX-IL2, remain tumor free after 7 weeks. These data clearly demonstrate the importance of an initial high dose of IL-2 to prevent tumor establishment.

Table 12
Effect of IL-2 modified fibroblasts on tumor establishment and development.
2 X 10⁶ fibroblasts mixed with 1 X 10⁵ CT26 tumor cells at time of injection.

Fibroblasts mixed with tumor cells	Animal Number	Tumor- bearing	Percent Tumor-free	Tumor Size (mm ²)			Mean Tumor Size (mm ²)		
				Total	25-100	101-200	201-300	>301	
After 6 Weeks*									
Control (no fibroblasts)**	13	0	13	0%	0	5	2	5	315 ± 197
Unmodified fibroblasts**	20	0	20	0%	0	2	3	14	350 ± 100
DCTK-IL2 fibroblasts	12	4	8	33%	0	1	4	3	185 ± 141
LXSN-IL2 fibroblasts***	15	4	11	27%	0	5	1	2	135 ± 121
LNCX-IL2 fibroblasts	8	6	2	75%	2	0	0	0	8 ± 14

* Mean tumor size is for 4 weeks, the last timepoint at which tumors were measured.

** One mouse in each of these arms developed an intraperitoneal tumor which was not measurable.

*** Three mice in this arm developed intraperitoneal tumors which were not measurable.

As an additional control, mice were injected with CT26 cells genetically modified to express IL-2 (results not shown). Injection of up to 1×10^6 IL-2 expressing tumor cells into Balb/c mice failed to produce tumors.

5 Injection of higher numbers however, resulted in some animals developing tumors with delayed onset. These data confirm the results reported in the literature (1). In order to compare the efficacy of IL-2 producing fibroblasts to IL-2 producing tumor cells, we mixed 2×10^6 CT26 tumor

10 cells modified with the DCTK-IL2 vector with 1×10^5 unmodified tumor cells. Figures 10 and 11, and Table 13 show that DCTK-IL2 modified tumor cells are somewhat effective in preventing tumor development. Four weeks after injection, the mean tumor size for the treatment arm

15 is 303 mm^2 , compared to 620 mm^2 for the control arm. After 22 weeks, one animal (10%) remains tumor free, compared to none in the control arms. Data for animals treated under the same conditions with DCTK-IL2 modified fibroblasts in a separate experiment are included for comparison purposes.

20 This comparison suggests that DCTK-IL2 modified tumor cells have an effect on tumor establishment similar to that of DCTK-IL2 modified fibroblasts.

Table 13

Effect of IL-2 modified cells on tumor establishment and development.
 2 X 10⁶ DCTK-IL2-modified CT26 tumor cells mixed with 1 X 10⁵ CT26 cells compared to 2 X 10⁶ DCTK-IL2-modified fibroblasts mixed with 1 X 10⁵ CT26.

Cells mixed with tumor cells	Animal Number			Tumor Size (mm ²)				Mean Tumor Size (mm ²)
	Total	Tumor- free	Tumor- bearing	Percent Tumor-free	25-100	101-200	201-300	
After 22 Weeks*								
Control (no fibroblasts)	5	0	5	0%	0	0	0	5
Unmodified fibroblasts	5	0	5	0%	0	0	0	5
DCTK-IL2-modified CT26 cells	10	1	9	10%	1	0	2	5
DCTK-IL2-modified fibroblasts	8	2	6	25%	0	1	2	3
								214 ± 158

* Mean tumor size is for 4 weeks, the last timepoint at which tumors were measured.

EXAMPLE VII(b)EFFECT OF FIBROBLAST MEDIATE CYTOKINE GENE THERAPY
ON SYSTEMIC ANTI-TUMOR IMMUNITY

Groups of Balb/c mice were immunized with
5 2.5×10^3 irradiated tumor cells either alone or mixed with
2 $\times 10^4$ transduced or unmodified fibroblasts, and challenged
one week later with 5×10^4 live tumor cells in the opposite
flank. These results (Figures 12 and 13, and Table 14)
demonstrate that immunization with irradiated tumor cells
10 and transduced fibroblasts protect some animals against a
live tumor challenge, but that the protection is only
slightly better than that achieved by immunization with
irradiated tumor cells alone or irradiated tumor cells
mixed with unmodified fibroblasts.

Table 14

Mice immunized with 2×10^6 fibroblasts mixed with 2.5×10^6 irradiated CT26 tumor cells 7 days prior to challenge with 5×10^4 fresh tumor cells.

Fibroblasts mixed with irradiated tumor cells	Animal Number		Percent Tumor-free	Tumor Size (mm ²)			Mean Tumor Size (mm ²)
	Total	Tumor-free		25-100	101-200	>301	
After 22 Weeks*							
Control (saline)	20	0	0%	0	0	1	19
Irradiated CT26 only**	16	5	31%	2	1	2	5
Irradiated CT26 mixed with unmodified fibroblasts	15	4	11	27%	0	1	3
DCTK-IL2 fibroblasts**	25	10	15	40%	4	1	8
							172 ± 194

* Mean tumor size is for 4 weeks, the last timepoint at which tumors were measured.

** One mouse in each of these arms developed an intraperitoneal tumor which was not measurable.

In a second protocol similar to the one described above, animals were challenged with fresh tumor cells two weeks following immunization with irradiated tumor cells mixed with fibroblasts. The results, shown in Figures 14 5 and 15, and in Table 15, demonstrate that DCTK-IL2 modified fibroblasts mixed with irradiated tumor cells confers superior protection to subsequent tumor challenge than irradiated tumor cells alone, irradiated tumor cells mixed with unmodified fibroblasts, or irradiated tumor cells 10 mixed with LNCK-modified fibroblasts. After 7 weeks, seven of ten animals (70%) treated with DCTK-IL2 modified fibroblasts remain tumor free compared to only one third of the control animals. At four weeks, the mean tumor size of this group was 41 mm³, compared to 180, 170, and 140 mm³ for 15 the three control groups. Animals treated with LNCK-IL2 modified fibroblasts were also protected against subsequent tumor challenge, but the results were less striking. In this group, 54% of the animals remain tumor free and the mean tumor size for the group at four weeks was 86 mm³. The 20 number of tumor free animals in the group treated with LXSN-IL2 modified fibroblasts was similar to the control groups, although the tumors were slightly delayed in their onset. A multivariate non-parametric statistical procedure (19, 20), utilized to evaluate differences in tumor onset, 25 demonstrated that the differences for the six arms presented in Figure 15 were significant (p = 0.012). It further showed that the saline control arm and the arms that received irradiated tumor cells alone or mixed with unmodified or LNCK vector modified fibroblasts formed a 30 statistical group. A second, distinct statistical group was formed by the three arms that received IL-2 vector modified fibroblasts mixed with irradiated tumor cells. Subsequent comparisons between the saline injected control 35 arm and animals that received tumor cells mixed with IL2 transduced fibroblasts revealed a significant difference for all vectors (p < 0.05).

Table 15

Mice immunized with 2×10^6 fibroblasts on induction of systemic anti-tumor immunity. Mice immunized with 2×10^6 fibroblasts mixed with 2.5×10^5 irradiated CT26 tumor cells 14 days prior to challenge with 5×10^4 fresh tumor cells.

Immunization by fibroblasts mixed with irradiated tumor cells	Animal Number	Tumor Size (mm ²)			Mean Tumor Size (mm ²)
		Total free bearing	Tumor-free bearing	Percent Tumor-free	
After 7 Weeks: ^a		25-100	101-200	201-300	>301
Control (alive)**	8	1	7	13%	0
Irradiated CT26 only	10	3	7	30%	0
Irradiated CT26 mixed with unmodified fibroblasts	6	2	4	33%	0
Irradiated CT26 mixed with LNCK-modified fibroblasts	10	3	7	30%	3
Irradiated CT26 mixed with LNCK-IL2-modified fibroblasts	13	7	6	54%	1
Irradiated CT26 mixed with LMSN-IL2-modified fibroblasts	12	4	8	33%	5
Irradiated CT26 mixed with DCTK-IL2-modified fibroblasts	10	7	3	70%	1

^a Mean tumor size is for 4 weeks, the last timepoint at which tumors were measured.

^{**} One mouse in this arm developed an intraperitoneal tumor which was not measurable.

These results demonstrate the feasibility of using genetically modified fibroblasts as a means of delivering cytokine gene therapy. In all experiments, the LNCX-L2 vector proved superior in preventing tumor establishment while the DCTK-IL2 vector was better in the induction of systemic protection against subsequent tumor challenges. These contrasting effects, although somewhat surprising, can be explained by the observation that the CMV promoter is turned off in vivo five days after 10 implantation while the TK promoter remains active for a longer period of time. The implication of this finding is that to apply this method of gene therapy successfully we have to use promoters that result in high level, sustained expression of IL-2 in vivo in the transduced fibroblasts.

15 The data obtained from this research effort has important implications for all cytokines that have either direct or indirect anti-tumor effects. Furthermore, this data suggests that anti-tumor efficacy is IL-2 dose dependent. Hence, construction of vectors which result in 20 higher levels of cytokine secretion will be a significant advance toward the application of this method of gene therapy.

Reference numbers in parenthesis in the above examples correspond to the following list of references and 25 are incorporated herein by reference.

References

1. Gabrilove, J.L. et al., *Monogr. J. Natl. Cancer Inst.* 10:73-7 (1990).
- 5 2. Kelso, A., *Current Opinion in Immunology*, 2:215-25 (1989).
3. Borden, E.C. et al., *Cancer*, 65:800-14 (1990).
4. Rosenberg, S.A. et al., *Ann. Intern. Med.*, 108:853-864 (1988).
- 10 5. Lotze, M.T. et al., *JAMA*, 256:3117-3124 (1986).
6. Pizza, G. et al., *Lymphokine Research*, 1:45-8 (1988).
7. Sarna, G. et al., *Journal of Biological Response Modifiers*, 9:81-6 (1990).
- 15 8. Gandolfi, L. et al., *Hepato-Gastroenterology*, 36:352-6 (1989).
9. Bubenik, J. et al., *Immunol. Letters*, 19:279-82 (1988).
10. Bubenik et al., *Immunol. Letters*, 23:287-292
- 20 25 11. Fearon, E.R. et al., *Cell*, 60:387-403 (1990).
12. Gansbacher, B. et al., *J. Exp. Med.*, 172:1217-1224 (1990).
13. Watanabe, Y. et al., *Proc. Natl. Acad. Sci.*, 86:9456-9460 (1989).

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14. Tepper, R.I. et al., *Cell*, 57:503-512 (1989).
15. Kriegler, M., *Gene Transfer and Expression: A Laboratory Manual*, Stockton Press (1990).
16. Rosenberg, S.A. et al., *N. Eng. J. Med.*, 330 (1990).
17. Cornetta, K. et al., *Prog. Nucl. Acid Res. Mol. Biol.*, 36:311-22 (1989).
18. Hoover, H.C. et al., *Cancer Res.*, 44:1671-76 (1984).
19. Sobol et al. *New Eng. J. Med.* 316:1111-1117 (1987).
20. Li Xu, et al., *Virology*, 171:331-341 (1989).

Although the invention has been described with reference to the presently-preferred embodiment, it should be understood that various modifications can be made without departing from the spirit of the invention.
5 Accordingly, the invention is limited only by the following claims.

WE CLAIM:

1. A method of treating cancer in a patient comprising the stimulation of that patient's immune response against the cancer by immunizing said patient at a site other than an active tumor site with a formulation 5 comprising tumor antigens and CE cells genetically modified to express at least one cytokine gene product.
2. The method of claim 1 wherein tumor cells previously isolated from said patient provide the tumor antigens.
3. The method of claim 1 wherein the cytokine gene is selected from the group consisting of interleukin-1, interleukin-2, interleukin-3, interleukin-4, interleukin-5, interleukin-6, and gamma-interferon.
4. The method of claim 3 wherein one cytokine gene is interleukin-2.
5. The method of claim 1 wherein at least one cytokine gene is transferred into cells to generate CE cells by recombinant methods.
6. The method of claim 5 wherein the cytokine gene is present in an expression vector.
7. The method of claim 6 wherein said expression vector additional contains a suicide gene.
8. The method of claim 5 wherein the CE cells are generated from fibroblasts and antigen-presenting cells.

9. A method for enhancing a patient's immune response to a cancer comprising:

- a) isolating fibroblasts from said patient;
- 5 b) culturing said fibroblasts *in vitro*;
- c) transducing said fibroblasts with a retroviral expression vector containing the gene coding for IL-2 and a gene coding for a tumor antigen in a retroviral expression vector, to express said tumor antigen and to express and secrete said IL-2 by said fibroblasts; and
- 10 d) immunizing said patient with said fibroblasts that express IL-2 at a level sufficient to enhance an immune response but low enough to avoid substantial systemic toxicity and that express said tumor antigen, at a site other than an active tumor site.
- 15
- 20

10. The method of claim 9 wherein said fibroblasts are further modified to express a suicide gene.

11. A composition for increasing a patient's immune response to tumor antigens comprising tumor antigens and CE cells genetically modified to express at least one cytokine gene product.

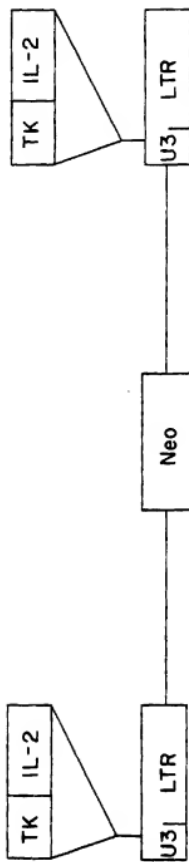
12. The composition of claim 11 wherein the cytokine gene is selected from the group consisting of interleukin-1, interleukin-2, interleukin-3, interleukin-4, interleukin-5, interleukin-6, and gamma interferon.

13. The composition of claim 12 wherein one cytokine gene is interleukin-2.

14. The composition of claim 11 wherein each cytokine gene is expressed at a level sufficient to stimulate the immune response but low enough to avoid substantial systemic toxicities.

15. The method of claim 9 wherein in said transducing step said retroviral expression vector has a promotor causing sustained secretion of IL-2.

16. The method of claim 15 wherein said retroviral expression vector causes the secretion of at least four units of IL-2 per day for a period of ten days or longer.



RETROVIRAL VECTOR LXSN-IL2



RETROVIRAL VECTOR LXSN-IL2



RETROVIRAL VECTOR LNCX-IL2

FIG. 1

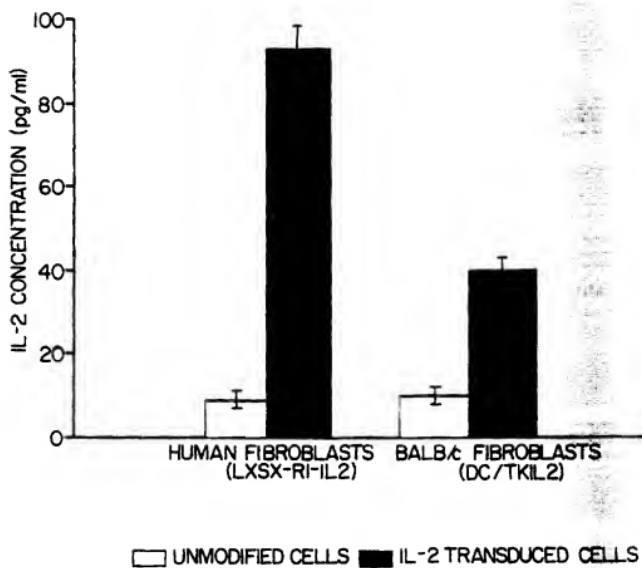


FIG. 2

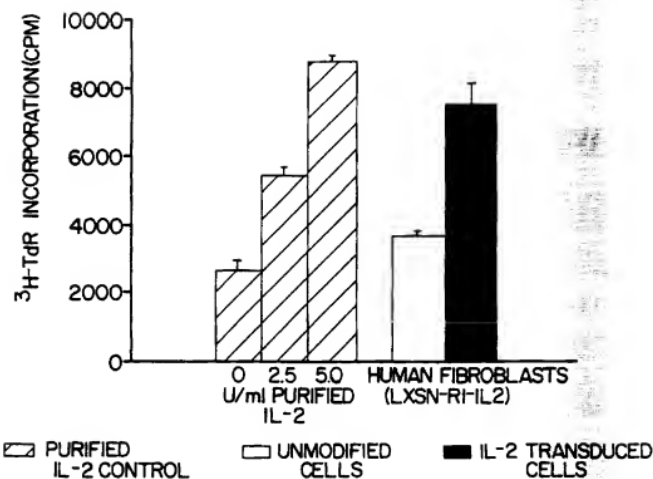


FIG. 3

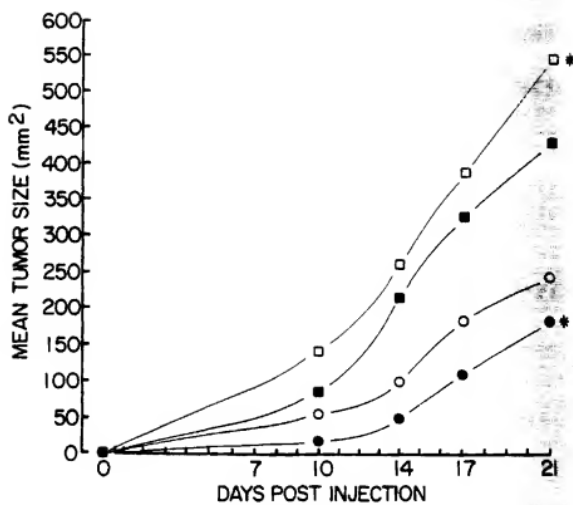


FIG. 4



1 2 3 4 5 6 7

FIG. 5

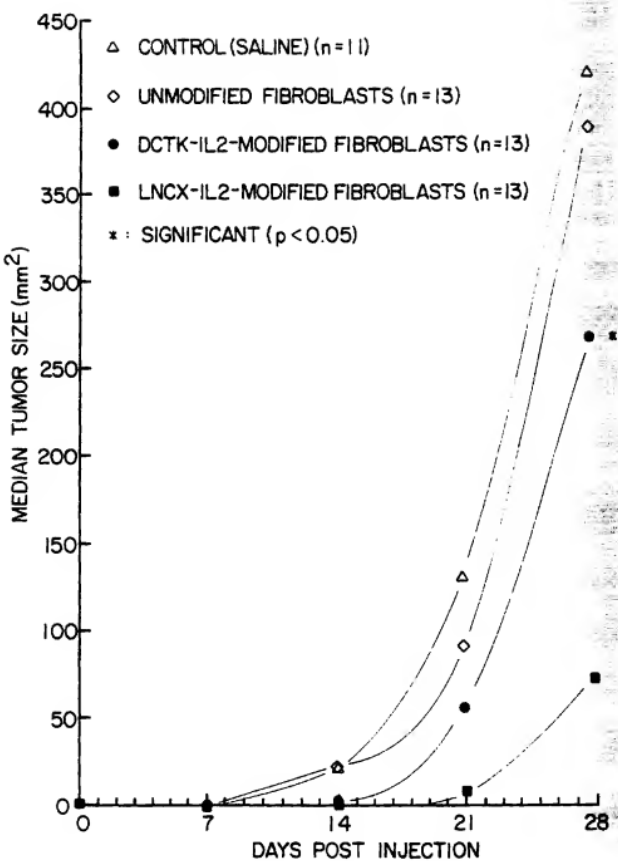
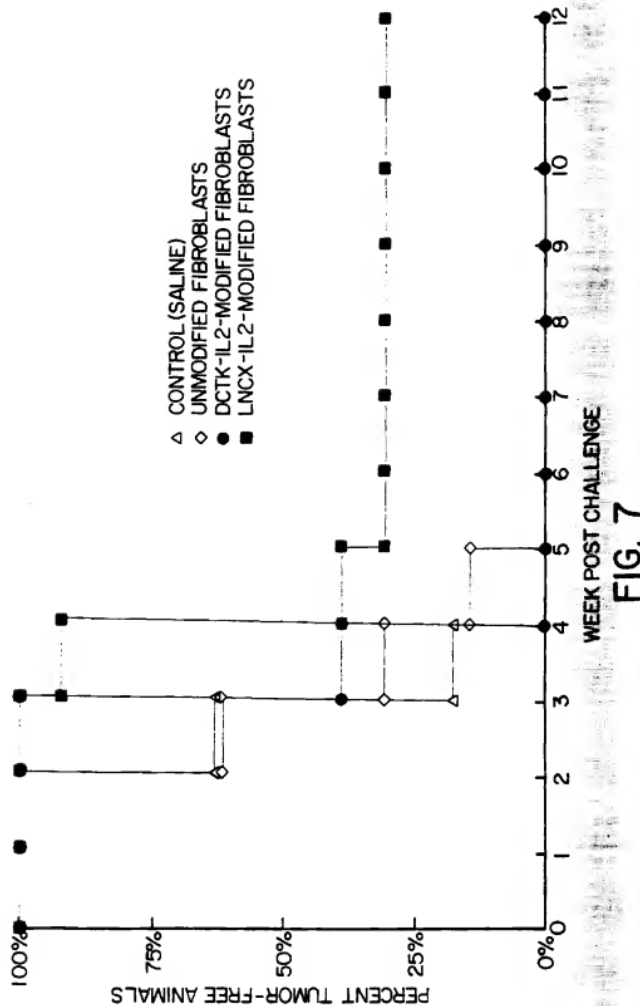


FIG. 6

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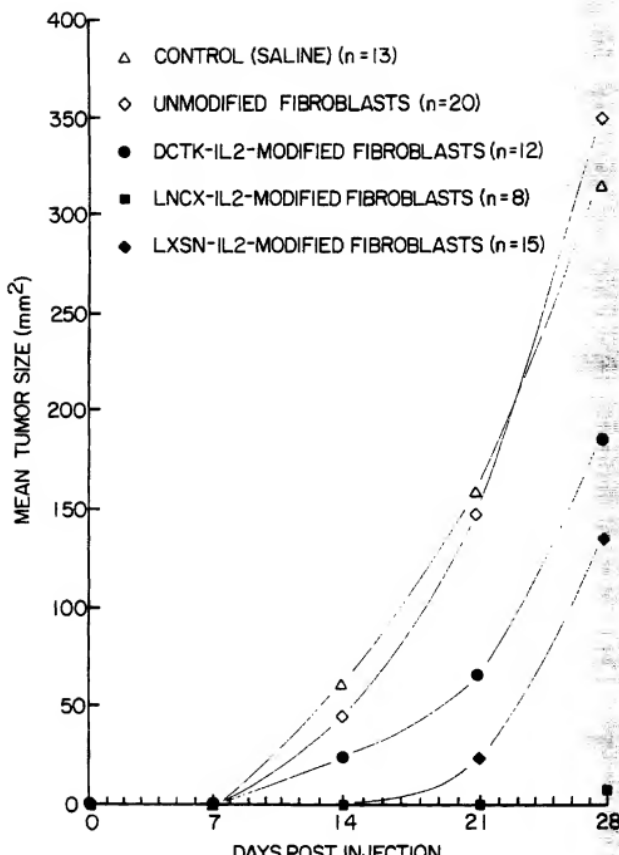


FIG. 8

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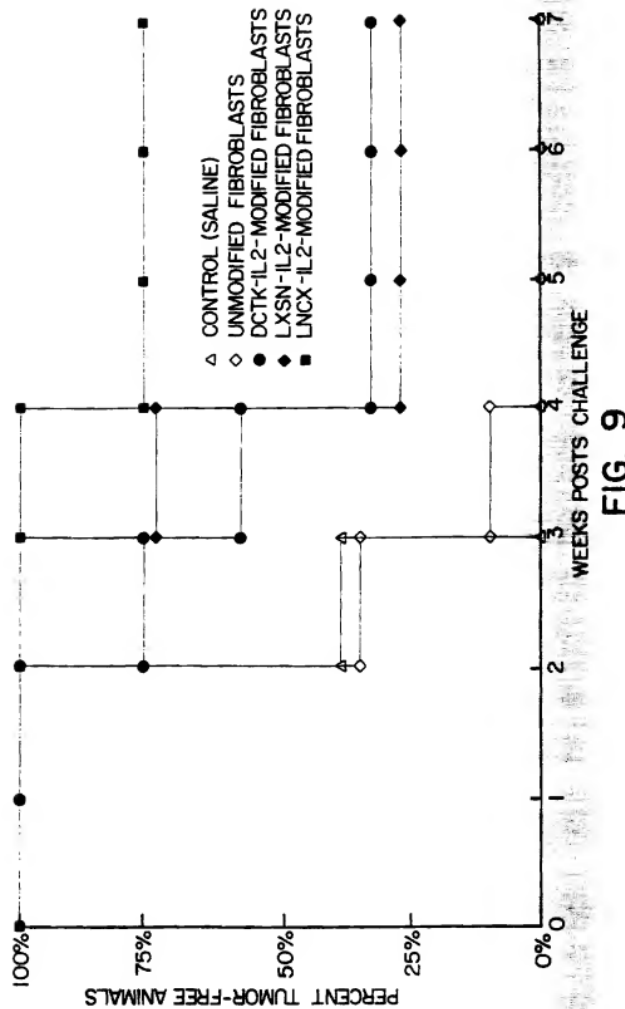


FIG. 9

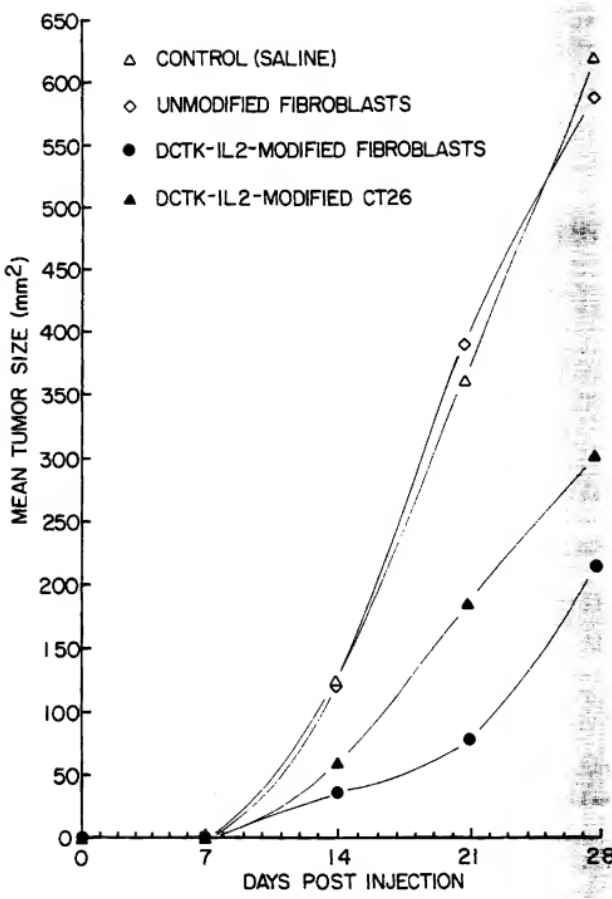


FIG. 10

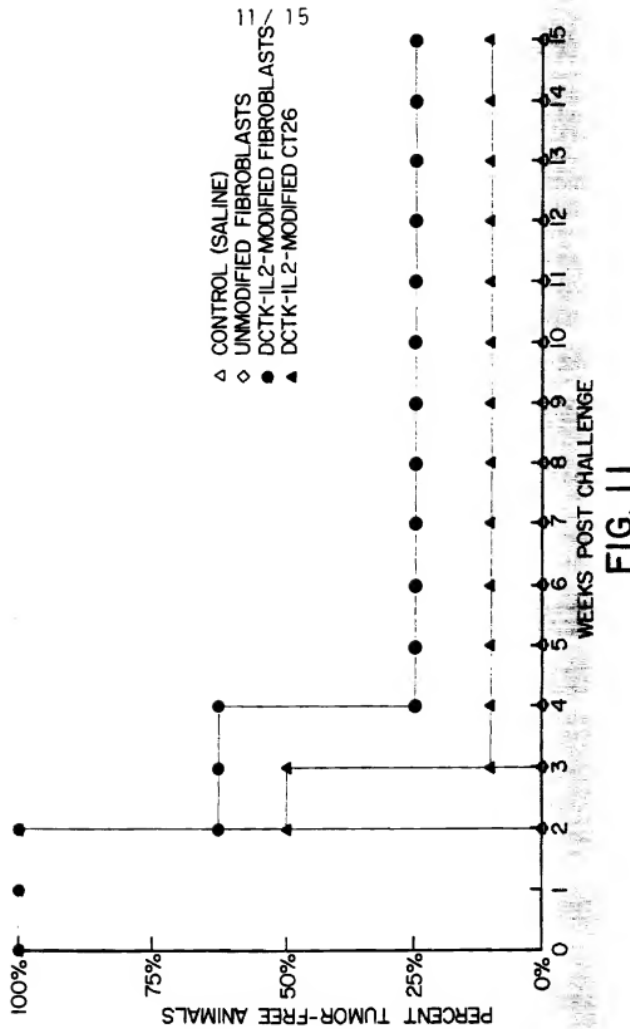


FIG. 11

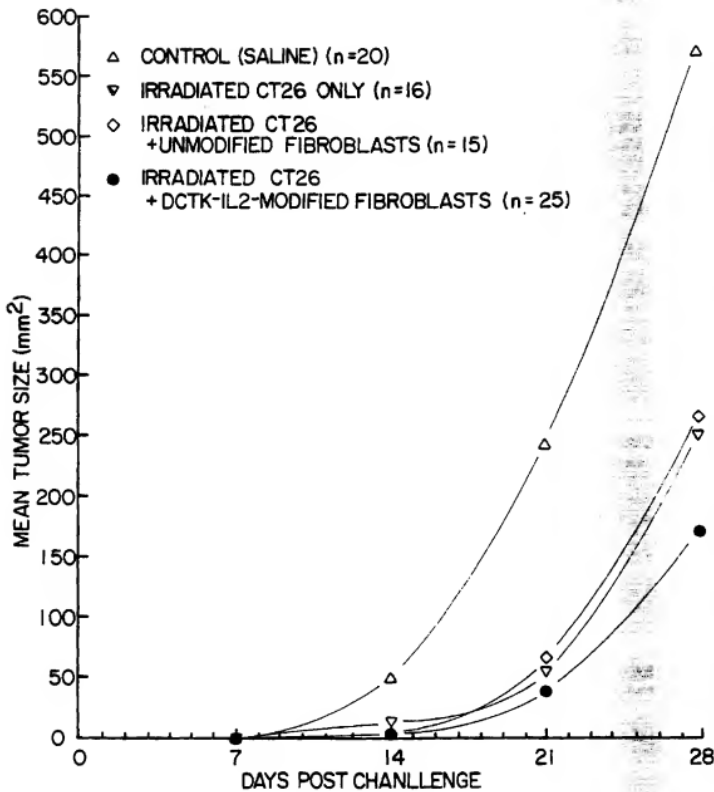
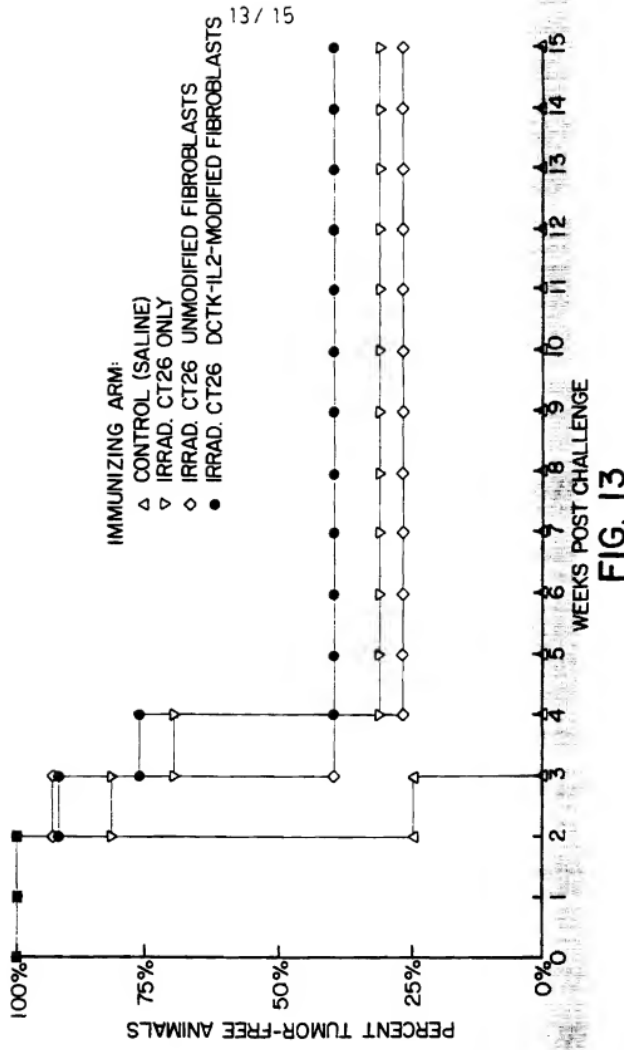


FIG. 12

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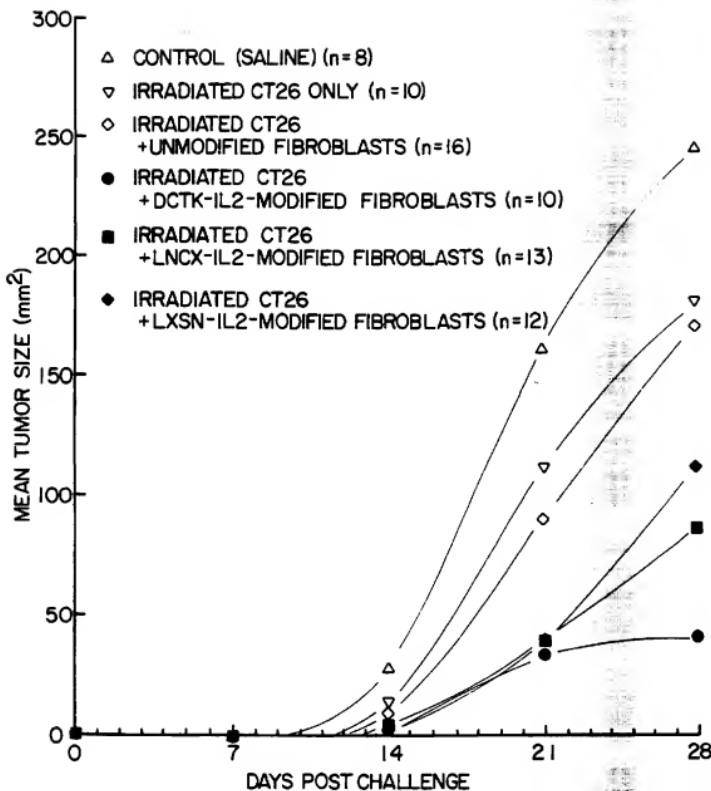
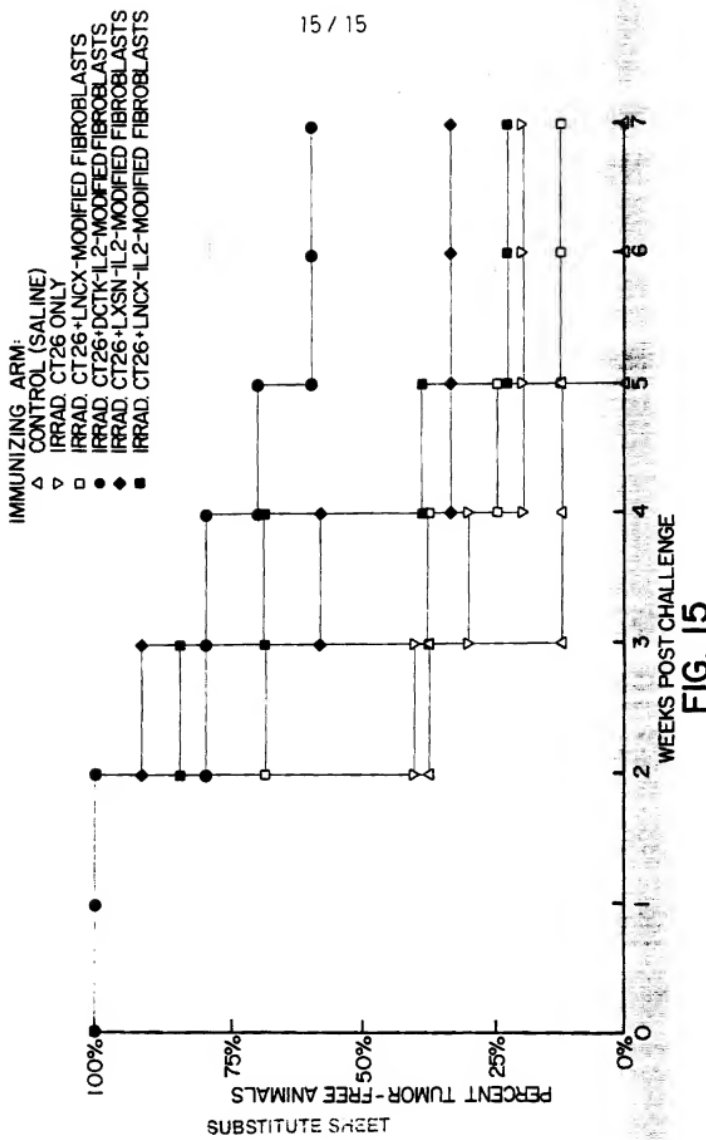


FIG. 14

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INTERNATIONAL SEARCH REPORT

International application No.

PCT/US92/08999

A. CLASSIFICATION OF SUBJECT MATTER

IPC(4) : Please See Extra Sheet.

US CL : Please See Extra Sheet.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 424/93B, 93U, 89; 435/240.2, 320.1, 69.5, 69.51, 69.52; 935/65, 32, 12, 57, 70, 71

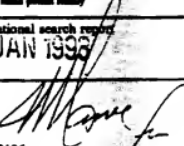
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

BIOSIS, MEDLINE, APS

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	Journal of Experimental Medicine, Volume 172, issued October 1990, Ganabacher et al., "Interleukin 2 Gene Transfer into Tumor Cells Abrogates Tumorigenicity and Induces Protective Immunity", pages 1217-1224, see the entire document.	1-8, 11-14 9, 10, 15, 16
Y	Cell, Volume 57, issued 05 May 1989, Tepper et al., "Murine Interleukin-4 Displays Potent Anti-Tumor Activity In Vivo", pages 503-512, see the entire document.	1-3, 5-6, 8-11, 12-14 4, 13
X	Cell, Volume 60, issued 09 February 1990, Fournier et al., "Interleukin-2 Production by Tumor Cells Bypasses T Helper Function in the Generation of an Antitumor Response", pages 397-403, see the entire document.	1-3, 5-8, 11-13 2, 6, 7, 14-16
Y	Cancer Research, Volume 50, issued 15 August 1990, Ogura et al., "Implantation of Genetically Manipulated Fibroblasts into Mice as Antitumor α -Interferon Therapy", pages 5102-5106, see the entire document.	1-16

<input checked="" type="checkbox"/>	Further documents are listed in the continuation of Box C.	<input type="checkbox"/>	See patent family annex.
* "A"	Special category of cited documents: document defining the present state of the art which is not considered to be part of particular relevance	"T"	later document published after the international filing date or priority date and not in conflict with the application but used to understand the principle or theory underlying the invention
"E"	earlier document published on or after the international filing date	"X"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"L"	document which may throw doubt on priority claim(s) or which is cited to establish the publication date of another citation or other special events (as specified)	"Y"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"O"	document referring to an oral disclosure, use, exhibition or other events	"A"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"P"	document published prior to the international filing date but later than the priority date claimed		document member of the same patent family
Date of the actual completion of the international search 11 January 1993	Date of mailing of the international search report 26 JAN 1993		
Name and mailing address of the ISA/ Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. NOT APPLICABLE	Authorized officer JACQUELINE STONE Telephone No. (703) 308-0196		

INTERNATIONAL SEARCH REPORT

Int. application No.

PCT/US92/08999

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	Cancer Research, Volume 50, issued 15 December 1990, Ganbacher et al., "Retroviral Vector-mediated Interferon Gene Transfer into Tumor Cells Generates Potent and Long Lasting Antitumor Immunity", pages 7820-7825, see the entire document.	1,3,5,6,8,11,12,14 2,7
Y		

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US92/08999

A. CLASSIFICATION OF SUBJECT MATTER:
IPC (5):

A61K 48/00, 35/12, 39/00; C12N 15/19, 15/24, 15/25, 15/26, 15/90, 15/63

A. CLASSIFICATION OF SUBJECT MATTER:
US CL :

424/93B, 93U, 89; 435/240.2, 320.1, 69.5, 69.51, 69.52; 935/65, 32, 12, 57, 70, 71